Connecting via Winsock to STN

STN Structure Search

10/526,507

Registry Caplus

Caplus

Welcome to STN International! Enter x:x

LOGINID: SSPTAJMN1626

PASSWORD:

NEWS 32

NEWS 33

NEWS 34

MAY 21

MAY 21

MAY 22

patents

patents

TERMINAL (ENTER 1, 2, 3, OR ?):2

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Web Page for STN Seminar Schedule - N. America
NEWS 2
       JAN 08
                CHEMLIST enhanced with New Zealand Inventory of Chemicals
                CA/CAplus Company Name Thesaurus enhanced and reloaded
NEWS 3 JAN 16
                IPC version 2007.01 thesaurus available on STN
NEWS 4
        JAN 16
NEWS 5
        JAN 16 WPIDS/WPINDEX/WPIX enhanced with IPC 8 reclassification data
NEWS 6 JAN 22
                CA/CAplus updated with revised CAS roles
NEWS 7
         JAN 22
                CA/CAplus enhanced with patent applications from India
NEWS 8
         JAN 29
                PHAR reloaded with new search and display fields
        JAN 29
NEWS 9
                CAS Registry Number crossover limit increased to 300,000 in
                 multiple databases
NEWS 10
         FEB 15
                 PATDPASPC enhanced with Drug Approval numbers
NEWS 11
         FEB 15
                 RUSSIAPAT enhanced with pre-1994 records
NEWS 12
         FEB 23
                 KOREAPAT enhanced with IPC 8 features and functionality
NEWS 13
        FEB 26
                MEDLINE reloaded with enhancements
NEWS 14
                 EMBASE enhanced with Clinical Trial Number field
         FEB 26
NEWS 15
         FEB 26
                 TOXCENTER enhanced with reloaded MEDLINE
NEWS 16
         FEB 26
                 IFICDB/IFIPAT/IFIUDB reloaded with enhancements
NEWS 17
         FEB 26
                CAS Registry Number crossover limit increased from 10,000
                 to 300,000 in multiple databases
NEWS 18
                 WPIDS/WPIX enhanced with new FRAGHITSTR display format
        MAR 15
NEWS 19
        MAR 16
                CASREACT coverage extended
NEWS 20
        MAR 20
                MARPAT now updated daily
NEWS 21
        MAR 22
                LWPI reloaded
NEWS 22
        MAR 30
                RDISCLOSURE reloaded with enhancements
NEWS 23
        APR 02
                 JICST-EPLUS removed from database clusters and STN
NEWS 24
        APR 30
                 GENBANK reloaded and enhanced with Genome Project ID field
NEWS 25
        APR 30
                CHEMCATS enhanced with 1.2 million new records
                CA/CAplus enhanced with 1870-1889 U.S. patent records
NEWS 26
        APR 30
NEWS 27
         APR 30
                 INPADOC replaced by INPADOCDB on STN
        MAY 01
NEWS 28
                 New CAS web site launched
NEWS 29
        MAY 08
                 CA/CAplus Indian patent publication number format defined
                 RDISCLOSURE on STN Easy enhanced with new search and display
NEWS 30
        MAY 14
                 fields
NEWS 31
        MAY 21
                 BIOSIS reloaded and enhanced with archival data
```

Welcome to STN International

NEWS EXPRESS NOVEMBER 10 CURRENT WINDOWS VERSION IS V8.01c, CURRENT MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP), AND CURRENT DISCOVER FILE IS DATED 25 SEPTEMBER 2006.

TOXCENTER enhanced with BIOSIS reload

CA/CAplus enhanced with additional kind codes for German

CA/CAplus enhanced with IPC reclassification in Japanese

10/526,507 05/25/2007

NEWS HOURS STN Operating Hours Plus Help Desk Availability

NEWS LOGIN Welcome Banner and News Items

NEWS IPC8 For general information regarding STN implementation of IPC 8

Enter NEWS followed by the item number or name to see news on that specific topic.

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FILE 'HOME' ENTERED AT 14:07:58 ON 25 MAY 2007

=> fil reg
COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 0.21 0.21

FULL ESTIMATED COST

FILE 'REGISTRY' ENTERED AT 14:08:03 ON 25 MAY 2007 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2007 American Chemical Society (ACS)

Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 24 MAY 2007 HIGHEST RN 935837-89-1 DICTIONARY FILE UPDATES: 24 MAY 2007 HIGHEST RN 935837-89-1

New CAS Information Use Policies, enter HELP USAGETERMS for details.

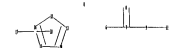
TSCA INFORMATION NOW CURRENT THROUGH December 2, 2006

Please note that search-term pricing does apply when conducting SmartSELECT searches.

REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/support/stngen/stndoc/properties.html

Uploading C:\Program Files\Stnexp\Queries\10526507\4.str



```
chain nodes :
4  6  7  8  9  11  17
ring nodes :
12  13  14  15  16
chain bonds :
6-7  7-8  7-9  9-11
ring bonds :
12-13  12-16  13-14  14-15  15-16
exact/norm bonds :
6-7  7-8  7-9  9-11  12-13  12-16  13-14  14-15  15-16
isolated ring systems :
containing 12 :
```

G1:Cb,Ak

G2:C,O,S,N

G3:C,N

G4:0,S

G5:H,Cb,Ak

Match level :

4:Atom 6:CLASS 7:CLASS 8:CLASS 9:CLASS 11:CLASS 12:Atom 13:Atom 14:CLASS

15:Atom 16:Atom 17:Atom 18:CLASS

Generic attributes :

4:

Saturation : Unsaturated

17:

Saturation : Unsaturated

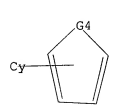
L1 STRUCTURE UPLOADED

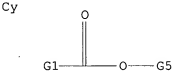
=> d

L1 HAS NO ANSWERS

Ll

STR





G1 Cb, Ak

G2 C, O, S, N

G3 C, N

G4 0, S

G5 H, Cb, Ak

Structure attributes must be viewed using STN Express query preparation.

=> s 11

SAMPLE SEARCH INITIATED 14:08:20 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 33022 TO ITERATE

6.1% PROCESSED

2000 ITERATIONS

50 ANSWERS

INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS:

ONLINE **COMPLETE**

BATCH **COMPLETE**

PROJECTED ITERATIONS:

649577 TO 671303

PROJECTED ANSWERS:

17610 TO 21354

L2

50 SEA SSS SAM L1

=> s 11 full

FULL SEARCH INITIATED 14:08:26 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 655775 TO ITERATE

100.0% PROCESSED 655775 ITERATIONS

19277 ANSWERS

SEARCH TIME: 00.00.10

L3 19277 SEA SSS FUL L1

=> save 13 apple/a ANSWER SET L3 HAS BEEN SAVED AS 'APPLE/A'

=> fil caplus
COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 172.55 172.76

FULL ESTIMATED COST

FILE 'CAPLUS' ENTERED AT 14:09:29 ON 25 MAY 2007
USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
PLEASE SEE "HELP USAGETERMS" FOR DETAILS.
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FILE COVERS 1907 - 25 May 2007 VOL 146 ISS 23 FILE LAST UPDATED: 24 May 2007 (20070524/ED)

Effective October 17, 2005, revised CAS Information Use Policies apply. They are available for your review at:

http://www.cas.org/infopolicy.html

=> s 13

L4 2272 L3

=>

Uploading C:\Program Files\Stnexp\Queries\10526507\4 A is thiazole.str



```
chain nodes :
5  6  7  8  10  16
ring nodes :
11  12  13  14  15  19  20  21  22  23
chain bonds :
5-6  6-7  6-8  8-10
ring bonds :
11-12  11-15  12-13  13-14  14-15  19-20  19-23  20-21  21-22  22-23
exact/norm bonds :
5-6  6-7  6-8  8-10  11-12  11-15  12-13  13-14  14-15  19-20  19-23  20-21  21-22
22-23
isolated ring systems :
containing 11 :
```

G1:Cb,Ak

G2:C,O,S,N

G3:C,N

G4:0,S

G5:H,Cb,Ak

Match level:
5:CLASS 6:CLASS 7:CLASS 8:CLASS 10:CLASS 11:Atom 12:Atom 13:CLASS 14:Atom 15:Atom 16:Atom 17:CLASS 19:Atom 20:Atom 21:Atom 22:Atom 23:Atom Generic attributes:
16:
Saturation : Unsaturated

L5 STRUCTURE UPLOADED

=> s 15 full sub=13
 REG1stRY INITIATED
Substance data SEARCH and crossover from CAS REGISTRY in progress...
Use DISPLAY HITSTR (or FHITSTR) to directly view retrieved structures.

FULL SUBSET SEARCH INITIATED 14:10:49 FILE 'REGISTRY'
FULL SUBSET SCREEN SEARCH COMPLETED - 1838 TO ITERATE

100.0% PROCESSED 1838 ITERATIONS SEARCH TIME: 00.00.01

. 400 ANSWERS

L6 400 SEA SUB=L3 SSS FUL L5

SUBSET IS IGNORED AS A SCOPE FOR THIS SEARCH L7 38 L6

=> s 17 and ppar 9158 PPAR L8 1 L7 AND PPAR

=> d ibib

```
L8 ANSWER 1 OF 1
ACCESSION NUMBER:
DOCUMENT NUMBER:
INVENTOR(S):

PATENT ASSIGNEE(S):
SOURCE:

PATENT ASSIGNEE(S):
SOURCE:

PATENT TYPE:
LINGUAGE:
FANILY ACC. NUM. COUNT:
PATENT NO.

PAT
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L8 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

CO. CR. CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KE, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MK, MZ, NI, NO, NZ, OM, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VC, VN, YU, ZA, ZM, ZW

RW: GH, GM, KE, LS, MM, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, EE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NI, PT, RO, SE, SI, SK, TR, BF, BT, CF, CG, CT, CM, GA, GQ, GW, MM, MR, NE, SN, TD, TG

AU 2003223233 A1 20030922 AU 2003-223233 20030307

AU 2003225692 A2 200309922 AU 2003-225662 20030307

AU 2003225692 A1 20030922 AU 2003-225662 20030307

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK, EP 1487443 A1 20041222 EP 2003-719363 20030307

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK, EP 1487443 A1 20041222 EP 2003-719363 20030307

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK, EP 1487443 A1 20041222 EP 2003-719363 20030307

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK, EP 1487443 A1 20050209 BR 2003-82184 20030307

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK, EP 1005525371 T 20050803 CN 2003-82184 20030307

JP 2005525371 T 20050803 CN 2003-82184 20030307

JP 2005525371 T 200501013 JP 2003-574149 20030307

JP 2005525371 T 200501013 JP 2003-574149 20030307

JP 2005525371 T 200501013 JP 2003-574149 20030307

JP 2005525075 T 20051013 JP 2003-574199 20060282

US 2006160796 A1 20060720 US 2006-385204 20060282

PRIORITY APPLN. INFO: US 2007-362702P P 200203088
                                                                                                                                                                                                                                                                                      US 2002-362702P
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                                                                                                                                                                                                                                                                                      WO 2003-US6784
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                                                                                                                                                                                                                                                                                      WO 2003-US7240
                                                                                                                                                                                                                                                                                                                                                                                                                  W 20030307
       OTHER SOURCE(S):
                                                                                                                                                                 MARPAT 139:6866
                                                                                                                                                                                                      THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE
       REFERENCE COUNT:
       FORMAT
```

=> d hitstr

- L8 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2007 ACS on STN

 IT 522440-69-0P
 RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
 (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
 (Vaes)

- (Uses)
 (antidiabetic and/or antitumor agent; preparation
 (benzylidene)rhodanines
 and analogs for treatment of cancer, diabetes, and other diseases)
 RN 532440-69-0 CAPUJS
 CN 3-Thiazolidineacetic acid, 5-[{5-(6-methoxy[1,1'-biphenyl]-3-yl)-2-furanyl]methylene]-4-oxo-2-thioxo-(9CI) (CA INDEX NAME)

=> d his

(FILE 'HOME' ENTERED AT 14:07:58 ON 25 MAY 2007)

FILE 'REGISTRY' ENTERED AT 14:08:03 ON 25 MAY 2007

L1 STRUCTURE UPLOADED

L2 50 S L1

L3 19277 S L1 FULL

SAVE L3 APPLE/A

FILE 'CAPLUS' ENTERED AT 14:09:29 ON 25 MAY 2007

L4 2272 S L3

L5 STRUCTURE UPLOADED

S L5

FILE 'REGISTRY' ENTERED AT 14:10:49 ON 25 MAY 2007

L6 400 S L5 FULL SUB=L3

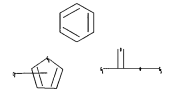
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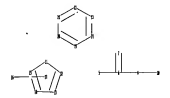
L7 38 S L6 SUBSET=L3 FULL

L8 1 S L7 AND PPAR

=>

Uploading C:\Program Files\Stnexp\Queries\10526507\4 A is phenyl.str





chain nodes::

5 6 7 8 10 16

ring nodes :

11 12 13 14 15 19 20 21 22 23 24

chain bonds :

5-6 6-7 6-8 8-10

10/526,507 05/25/2007

ring bonds :

 $11-\bar{1}2$ 11-15 12-13 13-14 14-15 19-20 19-24 20-21 21-22 22-23 23-24

exact/norm bonds :

5-6 6-7 6-8 8-10 11-12 11-15

normalized bonds :

12-13 13-14 14-15 19-20 19-24 20-21 21-22 22-23 23-24

isolated ring systems :

containing 11 :

G1:Cb,Ak

G2:C,O,S,N

G3:C, N

G4:0,S

G5:H,Cb,Ak

Match level :

5:CLASS 6:CLASS 7:CLASS 8:CLASS 10:CLASS 11:Atom 12:Atom 13:CLASS 14:Atom 15:Atom 16:Atom 17:CLASS 19:CLASS 20:Atom 21:Atom 22:Atom 23:Atom 24:Atom

Generic attributes :

16:

Saturation

: Unsaturated

L9 STRUCTURE UPLOADED

=> s 19 full sub=L3

REG1stRY INITIATED

Substance data SEARCH and crossover from CAS REGISTRY in progress... Use DISPLAY HITSTR (or FHITSTR) to directly view retrieved structures.

FULL SUBSET SEARCH INITIATED 14:13:34 FILE 'REGISTRY'
FULL SUBSET SCREEN SEARCH COMPLETED - 13743 TO ITERATE

100.0% PROCESSED 13743 ITERATIONS

0 ANSWERS

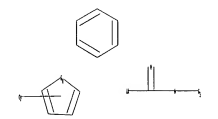
SEARCH TIME: 00.00.01

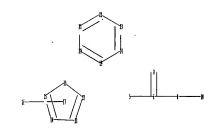
L10 0 SEA SUB=L3 SSS FUL L9

SUBSET IS IGNORED AS A SCOPE FOR THIS SEARCH L11 0 L10

=>

Uploading C:\Program Files\Stnexp\Queries\10526507\4 A is phenyl2.str





chain nodes :
5 6 7 8 10 16
ring nodes :
11 12 13 14 15 19 20 21 22 23 24
chain bonds :
5-6 6-7 6-8 8-10
ring bonds :
11-12 11-15 12-13 13-14 14-15 19-20 19-24 20-21 21-22 22-23 23-24
exact/norm bonds :
5-6 6-7 6-8 8-10 11-12 11-15
normalized bonds :
12-13 13-14 14-15 19-20 19-24 20-21 21-22 22-23 23-24
isolated ring systems :
containing 11 :

G1:Cb,Ak

G2:C,O,S,N

G3:C,N

G4:0,S

G5:H,Cb,Ak

Match level :

5:CLASS 6:CLASS 7:CLASS 8:CLASS 10:CLASS 11:Atom 12:Atom 13:CLASS 14:Atom 15:Atom 16:Atom 17:CLASS 19:CLASS 20:Atom 21:Atom 22:Atom 23:Atom 24:Atom

05/25/2007 10/526,507

Generic attributes :

16:

: Unsaturated Saturation

L12 STRUCTURE UPLOADED

=> s 112 full sub=L3 REG1stRY INITIATED

Substance data SEARCH and crossover from CAS REGISTRY in progress... Use DISPLAY HITSTR (or FHITSTR) to directly view retrieved structures.

FULL SUBSET SEARCH INITIATED 14:15:05 FILE 'REGISTRY' FULL SUBSET SCREEN SEARCH COMPLETED - 13754 TO ITERATE

100.0% PROCESSED 13754 ITERATIONS 0 ANSWERS

SEARCH TIME: 00.00.01

L13 O SEA SUB=L3 SSS FUL L12

SUBSET IS IGNORED AS A SCOPE FOR THIS SEARCH L140 L13

=> d his

L1

L4

(FILE 'HOME' ENTERED AT 14:07:58 ON 25 MAY 2007)

FILE 'REGISTRY' ENTERED AT 14:08:03 ON 25 MAY 2007

STRUCTURE UPLOADED

50 S L1 L2

19277 S L1 FULL L3

SAVE L3 APPLE/A

FILE 'CAPLUS' ENTERED AT 14:09:29 ON 25 MAY 2007

2272 S L3

L5 STRUCTURE UPLOADED

S L5

FILE 'REGISTRY' ENTERED AT 14:10:49 ON 25 MAY 2007

400 S L5 FULL SUB=L3 L6

FILE 'CAPLUS' ENTERED AT 14:10:50 ON 25 MAY 2007

38 S L6 SUBSET=L3 FULL

L7 1 S L7 AND PPAR L8

L9 STRUCTURE UPLOADED

S L9

FILE 'REGISTRY' ENTERED AT 14:13:33 ON 25 MAY 2007

L10 · 0 S L9 FULL SUB=L3

FILE 'CAPLUS' ENTERED AT 14:13:34 ON 25 MAY 2007

L11 0 S L10 SUBSET=L3 FULL

L12 STRUCTURE UPLOADED

S L12

FILE 'REGISTRY' ENTERED AT 14:15:05 ON 25 MAY 2007

L13 0 S L12 FULL SUB=L3

FILE 'CAPLUS' ENTERED AT 14:15:06 ON 25 MAY 2007

L14 0 S L13 SUBSET=L3 FULL

=> d ibib abs hitstr L7 1-38

L7 ANSWER 1 OF 38 CAPLUS COPYRIGHT 2007 ACS ON STN ACCESSION NUMBER: 2007:330181 CAPLUS DOCUMENT NUMBER: 146:358833 TITLE: PREPARATE ACCESSION NUMBER: 146:358833

146:358833
Preparation of thiazolinone and oxazolinone derivatives as PTP-1B inhibitors
Banerjee, Rakesh Kumar; Gupta, Ramesh Chandra; Tuli, Davinder; Rode, Milind; Shuthar, Bharat; Umrani, Dhananjay; Pathak, Padmaja; Choksi, Tejal; Chaudhary,

Anita Torrent Pharmaceuticals Ltd., India PCT Int. Appl., 110pp. CODEN: PIXXD2 PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: Patent

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

INVENTOR (S):

PAT	ENT	NO.			KIN	D	DATE			APPL	CAT	ION I	NO.		D	ATE	
						-									-		
WO	2007	0320	28		A1		2007	0322	1	WO 2	006-	IN36	8		2	0060	915
	W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,
		CN,	co,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,
		GE,	GH,	GM,	HN,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KM,	KN,	KP,
		KR,	KZ,	LA,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	LY,	MA,	MD,	MG,	MK,	MN,
		MW,	MX.	MY,	MZ,	NA,	NG,	NI,	NO.	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RS,
		RU,	sc,	SD,	SE,	SG,	SK,	SL,	SM,	sv,	SY.	TJ,	TM,	TN,	TR,	TT,	TZ.
		UA,	UG,	us,	UZ,	VC,	VN,	ZA,	ZM,	ZW							
	RW:	AT,	BE,	BG,	CH,	CY,	cz,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	IE,
		IS,	IT,	LT,	LU,	LV,	MC,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,	BJ,
		CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG,	BW,	GH,
		GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	AZ,	BY,
		KG,	KZ,	MD,	RU,	TJ,	TM										
PRIORITY	APP	LN.	INFO	.:						IN 2	005-	ко86	0		A 2	0050	916

OTHER SOURCE(S):

MARPAT 146:358833

The title thiazolinone and oxazolinone derivs. I [wherein ring A = naphthalene, biphenyl, etc.; ring B = (un)substituted (thiazolinone)methylene, (oxazolinone)methylene, etc.; ring C = benzene, naphthalene, etc.; L = NH, NHCH2, etc.; Y = (un)substituted CH2, CH2CH2, or CH2CH2CH2; R1 = H, -CH2CO2N, etc.; R2 and R3 = independently H,

ANSWER 1 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

REFERENCE COUNT:

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

ANSWER 1 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)
-CH2CO2H, etc.; R5 = COCO2H, (un) substituted CO2H, etc.; R8 and R9 =
independently H, halo, alkyl, etc.] or pharmaceutically acceptable salts
or prodrugs thereof are prepd. as protein tyrosine phosphatase (PPP)
inhibitors for treating or preventing PTP-1B mediated diseases. For
example, the compd. II was prepd. in a multi-step synthesis. Some of the
compds. I showed good inhibitory activities against human PTP-1B
92:9702-70-59 929703-11-79
BL: 28C (Pharmacological activity); SPM (Synthetic preparation). TMU Some of the

RE: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES

(drug candidate; preparation of thiazolinone and oxazolinone derivs.

85

PTP-1B inhibitors)
RN 929702-70-5 CAPIUS
CN Benzeneacetic acid,
4-[(5-[(2,2'-bithlophen]-5-ylmethylene)-4,5-dihydro-4oxo-2-thiazolyl]amino]-, sodium salt (1:1) (CA INDEX NAME)

929703-11-7 CAPLUS
Benzeneacetic acid, 4-[[4,5-dihydro-4-oxo-5-[(5-phenyl-2-thienyl)methylene]-2-thiazolyl]amino]-, sodium salt (1:1) (CA INDEX

L7 ANSWER 2 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
146:38337
Nolecular engineering of organic dyes containing
N-aryl carbazole moiety for solar cell
Kim, Duckhyun; Lee, Jae Kwan; Kang, Sang Ook; Ko,
Jaejung
CORPORATE SOURCE:
Department of Chemistry, Korea University, Chungnam,
339-700, S. Korea
SOURCE:
Tetrahedron (2007), 63(9), 1913-1922
CODEN: TETRAB: ISSN: 0040-4020
PUBLISHER:
Elsevier Ltd.
DOCUMENT TYPE:
JOURNAL
LANGUAGE:
Briglish
AB Organic dyes containing N-aryl carbazole moiety are designed and
synthesized.
Under standard global AM 1.5 solar condition, the JK-25 sensitized cell
gave a

Under standard grobal Am 1.5 south Constraints of 11.50 mA cm-2, an open circuit voltage (Voc) of 0.68 V, a fill factor of 0.66, corresponding to an overall conversion efficiency m of 5.15%, and the maximum incident monochromatic photon-to-current conversion efficiency (IPCE) of 77% at

nm. 930765-99-4P RL: PEP (Physical, engineering or chemical process); PRP (Properties);

(Purification or recovery); SPN (Synthetic preparation); TEM (Technical

or engineered material use); PREP (Preparation); PROC (Process); USES (Uses) (dye JK-26; mol. engineering of organic dyes containing N-aryl carbazole molety for solar cell)
RN 930765-99-4 CAPLUS
CN INDEX NAME NOT YET ASSIGNED

Double bond geometry as shown.

IT

930766-00-0P RL: PEP (Physical, engineering or chemical process); PRP (Properties);

(Purification or recovery); SPN (Synthetic preparation); TEM (Technical

INVENTOR (S): PATENT ASSIGNEE (S):

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

JP 2006337636

PRIORITY APPLN. INFO.:

PATENT NO.

SOURCE: DOCUMENT TYPE:

DATE

20050601

146:71778
Photographic materials containing cyanine dyes and silver halides sensitized with gold compounds Kataoka, Emiko
Konica Minolta Photo Imaging, Inc., Japan
Jpn. Kokai Tokkyo Koho, 72pp.
CODEN: JKXXAF

APPLICATION NO.

JP 2005-161101 JP 2005-161101

ANSWER 2 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued) engineered material use); PREP (Preparation); PROC (Process); USES (Uses) (dye JK-27; mol. engineering of org. dyes contg. N-aryl carbazole molety for solar cell) 930766-00-0 CAPLUS INDEX NAME NOT YET ASSIGNED

Double bond geometry as shown

REFERENCE COUNT:

34 THERE ARE 34 CITED REFERENCES AVAILABLE FOR

RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

L7 ANSWER 3 OF 38 CAPLUS COPYRIGHT 2007 ACS ON STN ACCESSION NUMBER: 2006:1309245 CAPLUS DOCUMENT NUMBER: 146:71778 Photographic materials contain

Patent

KIND

DATE

20061214

[L1Au(I)---(L2)m1]Mn1 II

AB Photog. materials having an emulsion layer containing cyanine dye I (Z1, Z2 =

groups for forming N-containing heterocycle: L1, L2, L3 = (un) substituted methine: A1, A2 = (un) substituted alkyl, aryl; V = thienyl; Y = counter ion; m = number for neutralization: n = 0-3; p, q = 0, 1) and 21 Ag halide particles sensitized with II (Au(I) = monovalent Au: L1 = groups bonding to Au(I) via N, S, O, P, Se, Te; L2 = compds. coordinating with Au(I) via N, S, O, P, Se, Te; M = H, counter cation; ml, nl = 0, 1}. The emulsions are stable and show excellent coating properties.

RE: MOA (Modifier or additive use); TEM (Technical or engineered material use); USES (USES)

(photog. materials with emulsion layers containing cyanine dyes and aold

compound-sensitized silver halides)
916613-49-5 CAPLUS
Benzothiazolium, 2-[[3-(carboxymethyl)-5-phenyl-2(3H)benzothiazolylidene|methyl]-3-(3-sulfopropyl)-5-(2-thienyl)-, inner salt,
sodium salt (1:1) (CA INDEX NAME)

ANSWER 3 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

L7 ANSWER 4 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 2006:1179845 CAPLUS
DOCUMENT NUMBER: 145:492279
TITLE: Semiconductors for photoelectric conversion, photoelectric conversion materials, and

dye-sensitized

solar cells
Kagawa, Nobuaki; Ofuku, Koji
Konica Minolta Holdings, Inc., Japan
Jpn. Kokal Tokkyo Koho, 60pp.
CODEN: JKCKARF
Patent
Japanese INVENTOR(S):
PATENT ASSIGNEE(S):
SOURCE:

Japanese

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. DATE APPLICATION NO. KIND DATE

JP 2006310097 PRIORITY APPLN. INFO.: JP 2005-131461 JP 2005-131461 20061109 20050428 20050428

OTHER SOURCE(S): MARPAT 145:492279

The title semiconductors have their surfaces adsorbing compds. I or II (ArI-2 = 5- or 6-membered aromatic ring, heterocycle; Hetl = bivalent heterocycle; X = 0, SOn; n = integer of 0-2, :NR; R = aliphatic, aryl, heterocycle; Cp = visible light- or near IR-absorbing group having 21 carboxyl; L1-5 = (un)substituted methine; ml, m2 = integer of 0-2). Further defined preferable Markush structures for I and II are

also
given. Also claimed are photoelec. conversion elements including the semiconductors and solar cells including the elements. The devices show high photoelec. energy conversion efficiency and are stable.

IT 914307-28-1 914307-34-9 914307-58-7 RL: DEV (Device component use); TEM (Technical or engineered material use); USES (Uses)
[methine dye-adsorbed semiconductors for solar cells)
RN 914307-28-1 CAPLUS
CN 3-Thiazolidineactic acid,
5-[[6-[5'-(butylethylamino][2,2'-bithiophen]-5-

(Continued)

ANSWER 4 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued) y11-4-[3-carboxy-1-{3,4-dichloropheny1}-1,5-dihydro-5-oxo-4H-pyrazol-4-ylidene]-4H-pyran-2-yl]methylene]-4-oxo-2-thioxo- (9CI) (CA INDEX NAME)

914307-34-9 CAPLUS 3-Thiazolidineacetic acid, 5-[2-[5-[7-(diethylamino)-2-oxo-2H-1-

benzopyran-3-yl]-2-thienyl]ethenyl]-6-[(1,3-diethyltetrahydro-2,4,6-trioxo-

5(2H)-pyrimidinylidene)methyl]-1,1-dioxido-4H-thiopyran-4-ylidene]-4-oxo-2-thioxo- (9CI) (CA INDEX NAME)

914307-58-7 CAPLUS 1-Piperidine Pigotion of the property of t

сн₂-- со₂н

ANSWER 4 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN INDEX NAME)

L7 ANSWER 5 OF 38 CAPLUS COPYRIGHT 2007 ACS ON STN ACCESSION NUMBER: 2006:681262 CAPLUS
DOCUMENT NUMBER: 145:145689

TITLE:

Preparation of rhodanine derivatives and analogs thereof as rigidified compounds for modulating heparanase activity Van-Gelder, Joel M.; Klein, Joseph Y.; Basel, Yochai; Reizelman, Anat; Tchilibon, Susanna; Mouallem, Orly Insight Blopharmaceuticals Ltd., Israel PCT Int. Appl., 193 pp. CODEN: PIXXD2
Patent INVENTOR (S):

PATENT ASSIGNEE (S):

DOCUMENT TYPE:

LANGUAGE: English

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

	TENT				KIN	D	DATE			APPL	ICAT	ION	NO.		_	ATE	
	2006				A2	-	2006	0713	1	WO 2	006-	1L23			_	0060	
WO	2006	0729	53		A3		2006	1102									
-	W:	AE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,
		CN,	co,	CR,	Cυ,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	ĸM,	KN,	ΚP,	KR,
		KZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	LY,	MA,	MD,	MG,	MK,	MN,	MW,	ΜX,
		MZ,	NA,	NG,	NI,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,
		SG,	sĸ,	SL,	SM,	SY,	TJ,	TM,	TN,	TR,	TT,	TZ,	UΑ,	υG,	US,	υz,	۷c,
		VN,	YU,	ZA,	ZM,	ZW											
	RW:	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	ΗU,	IE,
							MC,										
		CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	G₩,	ML,	MR,	ΝE,	SN,	TD,	TG,	BW,	GH,
		GM,	ΚE,	LS,	MW,	MZ,	NΑ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	ΑZ,	BY,
		KG,	ΚZ,	MD,	RU,	ΤJ,	TM										
PRIORIT	Y APP	LN.	INFO	.:					1	US 2	005-	6414	44P		P 2	0050	106

US 2005-681463P

P 20050517

GI

L7 ANSWER 5 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

Disclosed are novel rigidified compds. having a rhodanine-like residue

at least one aryl or heteroaryl residue linked to the rhodanine-like residue, whereby a core structure of these compds. I [Y and Z independently = O, S, Se, NRd, CRdRe or RdC=CRe; A = N or CRa; X = O, S, NRD, CRBRc etc.; M = N, P, C or Si; B = lone pair electrons, OR, alkoxy, amine, etc.; Ra, Rb, Rc, Rd, and Re independently = H, (un)substituted alkyl, cycloalkyl, and aryl; R1 = H, (un)substituted alkyl, cycloalkyl, alkenyl, etc.; core ring may be component of multicyclic system) is characterized as having one or zero free-to-rotate bonds. Methods for preparing I are included, e.g., II was prepared by cyclization of 2-(4-nitrophenyl)-4H-furo(3,2-b]pyrrole-5-carboxylic acid Et ester paraction

(preparation given) with 4-isothiocyanatobutyrate. In assays to determine inhibition

of heparanase (H53), I demonstrated IC50 values from 9.0-74.0 µM (with select compds. showing no inhibition). Also disclosed are pharmaceutical compns. containing these rigidified compds. and uses thereof for modulating the activity of heparanase and hence in the treatment of heparanase-associated diseases and disorders, and uses thereof for modulating the activity of heparin-binding proteins and hence in the treatment of heparin-binding proteins-associated diseases and disorders as well as in the

treatment of medical conditions that are at least partially treatable by rhodanine or a rhodanine analog.
898552-69-7P
RI: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES

(preparation of rhodanine derivs. and analogs thereof as rigidified

compound

ACCESSION NUMBER:

ACCESSION NUMBER:

DOCUMENT NUMBER:

145:159080

FURAN-2-ylmethylene Thiazolidinediones as Novel,
Potent, and Selective Inhibitors of Phospholnositide
3-Kinase y
Pomel, Vincent; Klicic, Jasna: Covini, David; Church,
Dennis D.; Shaw, Jeffrey P.; Roulin, Karen;
Burgat-Charvillon, Fablenne; Valognes, Delphine;
Camps, Montserrat; Chabert, Christian; Gillieron,
Corinne; Francon, Bernard, Perrin, Dominique, Leroy,
Didier; Gretener, Denise; Nichola, Anthony; Vitte,
Pierre Alain; Carboni, Susanna; Rommel, Christian;
Schwarz, Matthias K.; Rueckle, Thomas

Departments of Chemistry, Signal Transduction
Biochemical Pharmacology and Experimental
Pharmacology, Serono Pharmaceutical Research
Institute, Geneva, CH-1228, Switz.

Journal of Medicinal Chemistry (2006), 49(13),
3857-3871
CODEN: JNCMAR; ISSN: 0022-2623
American Chemical Society
Journal
DOCUMENT TYPE:
DOCUMENT TYPE:
Journal

PUBLISHER: DOCUMENT TYPE: LANGUAGE: GI

Class I phosphoinositide 3-kinases (PI3Ka), in particular PI3Ky, have become attractive drug targets for inflammatory and autoimmune diseases. Here, we disclose a novel series of furan-2-ylmethylene thiazolidinediones as selective, ATP-competitive PI3Ky inhibitors. Structure-based design and X-ray crystallog, of complexes formed by inhibitors bound to PI3Ky inheritate the properties of t

of acute peritonitis led to a significant reduction of leukocyte

L7 ANSWER 7 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2006:232172 CAPLUS DOCUMENT NUMBER: 144:295880 Semiconductor electrodes using

144:29580 Semiconductor electrodes using polymethine dye photosensitizers, photoelectric conversion devices, and dye-sensitized solar cells Fukui, Atsushi, Enomoto, Kazuhiro

INVENTOR (S): PATENT ASSIGNEE(S): SOURCE: Sharp Corp., Japan
Jpn. Kokai Tokkyo Koho, 21 pp.
CODEN: JKXXAF

DOCUMENT TYPE: Patent

Japanese

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

DATE APPLICATION NO. DATE PATENT NO. KIND JP 2004-256095 JP 2004-256095 20060316 JP 2006073375 PRIORITY APPLN. INFO.: А

OTHER SOURCE(S): MARPAT 144:295880

The title electrodes are equipped with semiconductors sensitized by polymethine dyes having bisenamine skeletons I (A and B = lower alkyl, (substituted) aryl, R = lower alkyl, (substituted) aryl or aralkyl; Rl-R4 = H, halogen, lower alkyl, or lower alkoxy; R and Rl or R and R3 may form 5- or 6-membered C rings; R1 and R2 or R3 and R4 may form 6-membered C rings; R5 to R7 = H, lower alkyl, or lower alkoxy; X = (substituted) aric

atic
group; Y = (substituted) heterocyclic group; n and m = 0-3 integer). The
resulting solar cells provide high photoelec. conversion efficiency.
879211-93-5
RL: DEV (Device component use); USES (Uses)
(dye; semiconductor electrodes using polymethine dye photosensitizers
for dye-sensitized solar cells)
879211-93-5 CAPLUS
3-Thiazolidineacetic acid, 5-[[5-[1-(2,2-diphenylethenyl)-1,2,3,4tetrahydro-6-quinolinyl]-2-thienyl]methylene]-4-oxo-2-thioxo- (9CI)
INDEX NAME)

ANSWER 6 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued) 900515-02-8P 900515-05-1P RL: PAC (Pharmacological activity); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation) (furan-2-ylmethylene thiazolidinediones as novel, potent, and ective inhibitors of phosphoinositide 3-kinase y) 900515-02-8 CAPLUS Benzeneacetic acid, 2-(5-[(2)-(2,4-dioxo-5-thiazolidinylidene)methyl)-2-furanyl]- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

900515-05-1 CAPLUS
Benzeneacetic acid, 3-[5-[(Z)-(2,4-dioxo-5-thiazolidinylidene)methyl]-2-furanyl]- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

REFERENCE COUNT:

THERE ARE 58 CITED REFERENCES AVAILABLE FOR 58

RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

ANSWER 7 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

L7 ANSWER 8 OF 38 CAPLUS COPYRIGHT 2007 ACS ON STN ACCESSION NUMBER: 2005:695895 CAPLUS
DOCUMENT NUMBER: 143:196809

DOCUMENT NUMBER: 143:195809
Coumarin dye-sensitized semiconductors, photoelectric conversion materials, and solar cells
Daifuku, Koji: Kagawa, Nobuaki
Konica Minolta Holdings, Inc., Japan
Jpn. Kokai Tokkyo Koho, 41 pp.
CODEN: JKXXAF TITLE:

INVENTOR (S): PATENT ASSIGNEE (S):

SOURCE: DOCUMENT TYPE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE JP 2005209359 PRIORITY APPLN. INFO.: 20050804 JP 2004-11512 JP 2004-11512 20040120 20040120

OTHER SOURCE(S):

MARPAT 143:196809

AB The semiconductors, preferably metal oxides or sulfides, contain I

(having

21 CO2M; M = H, cation; R1, R2 = H, alkyl, aryl, heterocyclic
group; R3-R9 = H, substituent; R1R2 and R3R4 may form ring; R, R101 =
substituent; X1 = O, S, NR21; X2 = O, S; X3 = O, S, heterocycle; R21 = H,
substituent; Z = nonmetal atomic group necessary for forming 5- or
6-membered
heterocycle; m, n = O-2). Solar cells using the semiconductors show high
photoelec. conversion efficiency and durability.

18 61907-91-7 861907-92-8 681908-04-5.
RL: DEV (Device component use); MOA (Modifier or additive use); TEM
(Technical or engineered material use); USES (Uses)
(coumarin dye-sensitized semiconductors for photoelec. conversion
materials and solar cells)

RN 861907-91-7 CAPLUS
CN 3-Thiazolidineacetic acid, 4-oxo-5-[[5'-(2,3,6,7-tetrahydro-1,1,7,7tetramethyl-11-oxo-1H,5H,11H-[1]benzopyrano[6,7,8-ij]quinolizin-10yl)[2,2'-bithiophen]-5-yl]methylene]-2-thioxo-(9CI) (CA INDEX NAME)

L7 ANSWER 9 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2005:638246 CAPLUS

DOCUMENT NUMBER:

COPYRIGHT 2007 ACS on SIN 2005-638246 CAPLUS 143:156316 Semiconductor electrodes sensitized with γ-pyrone polymethine dyes, and solar cells using them

INVENTOR (S): PATENT ASSIGNEE (S): SOURCE:

them Fukul, Atsushi; Han, Li-Yuan; Enomoto, Kazuhiro Sharp Corp., Japan Jpn. Kokai Tokkyo Koho, 21 pp. CODEN: JKXXAF Patent

DOCUMENT TYPE:

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 2005197115	А	20050721	JP 2004-3153	20040108
PRIORITY APPLN. INFO.:			JP 2004-3153	20040108

OTHER SOURCE(S):

MARPAT 143:156316

$$\begin{array}{c|c}
 & & L & COOM \\
 & & COOM \\
R^6 & R^5 & R^4 \\
 & & C = C \\
 & C$$

The electrodes have sensitizing dyes I [A = C4-5 heterocyclylene; Z = C6-24 bivalent aromatic group; R1 = H, C1-4 alkoxy; R2 = H, C1-4 alkyl,

C6-24 bivaient aromatic group: Ni = N, C1-4 alkoxy; NZ = N, C1-4 alkyl,
C1-4 alkoxy, C6-12 aryl, (substituted) styryl; R1R2 may form 5- or
6-membered carbon ring; R3-R6 = H, halo, C1-4 alkyl, C1-4 alkoxy; L = H,
electron withdrawing group; M = H, cation; m = 0-3; n = 0-4]. Thus, TiO2
electrods ensistized with polymethine dye having y-pyrone and
benzothiazole structure II is exemplified.
859829-02-0P
RL: DEV (Device component use); IMF (Industrial manufacture); NOA
(Modifier or additive use); PREP (Preparation): USES (Uses)
(semiconductor electrodes sensitized with y-pyrone polymethine
dyes for solar cells)
859829-02-0 CAPLUS
Acetic acid, cyano[2-ethyl-6-[2-[5'-[(3-hexyl-2(3H)benzothiazolylidene|methyl][2,2'-bithiophen]-5-yl]ethenyl]-4H-pyran-4ylidene)- (9CI) (CA INDEX NAME)

ANSWER 8 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

861907-92-8 CAPLUS
3-Thiazolidineacetic acid, 5-[3-(carboxymethyl)-4-oxo-5-[[5'-{2,3,6,7-tetrahydro-1,1,7,7-tetramethyl-11-oxo-1H,5H,1HH-[1]benzopyrano[6,7,8-i]]quinolizin-10-yl]{2,2'-bithiophen]-5-yl]methylene]-2-thiazolidinylidene]-4-oxo-2-thioxo- (9CI) (CA INDEX NAME)

861908-04-5 CAPLUS
1(2R)-Pyrimidinacetic acid, 2-[3-(carboxymethyl)-4-oxo-2-thioxo-5-thiazolidinylidene]-5-[[5'-[7-(dibutylamino)-2-oxo-2R-1-benzopyran-3-yl][2,2'-bithiophen]-5-yl]methylene]tetrahydro-4,6-dioxo-3-phenyl- (9CI)(CA INDEX NAME)

L7 ANSWER 9 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

L7 ANSWER 10 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 2005:409306 CAPLUS
1005:409306 CAPLUS
142:441839
Rhodanine compounds and compositions for use as antiviral agents
Rajinder, Singh; Usha, Ramesh; Clough, Jeffrey: Issakani, Sarkiz D.; Look, Gary Charles
Rigel Pharmaceuticals, Inc., USA
PCT Int. Appl., 82 pp.
CODEN: PIXXD2
DOCUMENT TYPE: PATENT INFORMATION:
English
FAMILV ACC. NUM. COUNT: English
PATENT INFORMATION: DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: 20041028 BZ, CA, CH, FI, GB, GD, KR, KZ, LC, MZ, NA, NI, SK, SL, SY, 2A, 2M, ZW ZM, ZW, AM, C2, DE, DK, PT, RO, SE, ML, MR, NE,

US 2003-526726P

P 20031203

R SOURCE(S): MARPAT 142:441839

The invention describes compds. and pharmaceutical compns. useful as inhibitors of the ubiquitination. The compds. and compns. of the invention are useful as inhibitors of the biochem. pathways of organisms in which ubiquitination is involved. In particular, the compds. and compns. are useful for treating diseases caused by viruses such as poxviruses and retroviruses. The invention further provides for methods of treating smallpox, herpes virus and HIV infection in patients using the compds. OTHER SOURCE(S):

compns. of the invention. Preparation of selected rhodanine compds. is described. 851305-41-4

851305-41-4
RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL
(Biological study); USES (Uses)
(Rhodanine compds. and compns. for use as antiviral agents)
851305-41-4 CAPLUS
Phenylalanine,
(52)-5-[[5-(2-chlorophenyl)-2-furanyl]methylene]-4-oxo2-thioxo-3-thiazolidinyl]acetyl}-, 1,1-dimethylethyl ester (9CI) (CA
INDEX NAME)

Double bond geometry as shown.

ANSWER 11 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN SSION NUMBER: 2005:57682 CAPLUS MENT NUMBER: 142:159516

ACCESSION NUMBER: DOCUMENT NUMBER:

The photoelectric conversion material, semiconductor electrode and photoelectric converter which uses the TITLE:

electrode and photoelectric converte electrode Horiuchi, Tamotsu; Maruyama, Atsushi Mitsubishi Paper Mills, Ltd., Japan Jpn. Kokai Tokkyo Koho, 19 pp. CODEN: JKXXAF Patent INVENTOR(S): PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

DATE PATENT NO. KIND DATE APPLICATION NO. JP 2005019251 PRIORITY APPLN. INFO.: А 20050120 JP 2003-183493 JP 2003-183493 20030626

OTHER SOURCE(S): MARPAT 142:159516

The material uses a compound I $\{R1 = (substituted) \ alkyl, (substituted) \ aralkyl, (substituted) \ alkenyl, (substituted) \ allyl, or (substituted) \ heterocyclic ring; <math>R2 = linking \ group \ forming \ a \ structure \ with N: R1 \ and R2 \ may form a ring; R2, N \ and bonded benzene ring may bond to form a$

R3 = H, halo, (substituted) alkyl, or (substituted) alkoxy group: R4 = bivalent aromatic condensed ring or bivalent heterocyclic ring; R5 = = substituent having acidic group). The electrode has a semiconductor

croated on a surface-conductive substrate and a pigment adsorbed on the semiconductor layer; where the pigment contains 21 above compound I. The converter, especially for a photoelectrochem. cell, uses the above electrodes.

1. פערטאי-14-7 RL: MOA (Modifier or additive use); USES (Uses) (semiconductor electrodes containing sensitizing pigments for photoelec.

colec. .
converters in photoelectrochem. cells)
829097-74-7 CAPLUS
3-Thiazolidineacetic acid, 5-[[5'-[4-[4-(2,2-diphenylethenyl)phenyl]1,2,3,3a,4,8b-hexahydrocyclopent[b]indol-7-yl][2,2'-bithlophen]-5yl]methylene]-4-oxo-2-thioxo-(9CI) (CA INDEX NAME)

L7 ANSWER 10 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

L7 ANSWER 11 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

(Continued)

(Continued)

THERE ARE 33 CITED REFERENCES AVAILABLE FOR RECORD. ALL CITATIONS AVAILABLE IN THE RE

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L7 ANSWER 12 OF 38 CAPLUS COPYRIGHT 2007 ACS ON STN ACCESSION NUMBER: 2004:1068070 CAPLUS DOCUMENT NUMBER: 142:168973 A Novel Class of Inhibitors of
                                                                                                                                                                                                         142:168973
A Novel Class of Inhibitors of Peptide Deformylase Discovered through High-Throughput Screening and Virtual Ligand Screening Howard, Michael H.: Cenizal, Teodorica; Gutteridge, Steven; Hanna, Wayne S.: Tao, Yong; Totrov, Maxim; Wittenbach, Vernon A.; Zheng, Ya-Jun Stine-Haskell Research Center, DuPont Crop
       AUTHOR (S):
CORPORATE SOURCE:

Stine-Haskell Research Center, DuPont Crop

SOURCE:

Journal of Medicinal Chemistry (2004), 47(27),
6669-6672

CODEN: JMCMAR; ISSN: 0022-2623

PUBLISHER:
American Chemical Society

JOURENT TYPE:
JOURENT TYPE:
JOURNAL CASHEACT 142:168973

ABP Peptide deformylase (PDF) has been identified as a promising

antibacterial
and herbicide target. A structurally novel class of inhibitors

containing a
2-thioxo-thiazolidin-4-one heterocycle substituted by an arylidene group
at the 5-position and a hexanoic acid side chain at the 3-position was
discovered independently via high-throughput screening and virtual ligand
screening. Data mining and analog synthesis established a
structure-activity relationship for the side chain region that is
consistent with the docked structure.

Janobe-48-3 370096-48-3 370096-55-2

RL: PAC (Pharmacological activity); PRP (Properties); THU (Therapeutic
use); BIOL (Biological study); USES (Uses)
(peptide deformylase inhibitors: high-throughput and virtual ligand
screening and preparation)

N 370096-48-3 CAPLUS

N 3-Thiazolidinehexanoic acid, 5-(3-(5-methyl-2-furanyl)-1-phenyl-1H-
pyrazol-4-yl]methylene]-4-oxo-2-thioxo- (9CI) (CA INDEX NAME)
       CORPORATE SOURCE:
Protection,
```

REFERENCE COUNT:

FORMAT

ANSWER 13 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN HO2C-CH2-CH2 676647-88-4 CAPLUS
3-Thiazolidinebutanoic acid, 4-oxo-5-([2,2':5',2''-terthiophen]-5ylmethylene)-2-thioxo- (9CI) (CA INDEX NAME)

ANSWER 12 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

370096-55-2 CAPLUS
3-Thiazolidinehexanoic acid,
--5-[[1-phenyl-3-(2-thienyl)-1H-pyrazol-4yl]methylene]-2-thioxo- (9CI) (CA INDEX NAME)

HO2C- (CH2)3

676654-51-6 CAPLUS CN 3-Thiazolidinebutanoic acid,
5-[(3,4-diphenyl-2-furanyl)methylene)-4-oxo-2thioxo- (9CI) (CA INDEX NAME)

L7 ANSWER 13 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2004:927010 CAPLUS DOCUMENT NUMBER: 141:376382 Pinl-modulating compounds and methods of use for the treatment of Pinl-associated diseases, including TITLE: cancer
Bao, Lere; Kimzey, Amy
pintex Pharmaceuticals, Inc., USA
PCT Int. Appl., 189 pp.
CODEN: PIXXD2
Patent INVENTOR (S): PATENT ASSIGNEE (S): SOURCE: DOCUMENT TYPE: LANGUAGE: Facent English FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. DATE 20041104 20060803 A2 A3 WO 2004093803 WO 2004093803 WO 2004-US11957 20040416

2004093803 A3 20060803

W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, II, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MM, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, DS, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZM, RW: BM, GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZM, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GM, GQ, GW, ML, MR, NE, SN, TD, TG PRIORITY APPLN. INFO.: US 2003-463271P P 20030416 OTHER SOURCE(S): MARPAT 141:376382

AB The invention is directed to modulators, e.g., inhibitors, of Pin1 and Pin1-related proteins and the use of such modulators for treatment of associated states, e.g., for the treatment of cancer. The present associated states, e.g., for the treatment of cancer. The present invention
aims to provide photochemotherspeutic compds. with increased specificity as compared with known agents.

IT 312601-59-4 676647-89-4 676534-51-6
RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
{Pinl-modulating compds. for treatment of Pinl-associated diseases, including cancer)
RN 312601-58-4 CAPLUS
CN 3-Thiazolidinepropanoic acid,
4-oxo-5-[[1-phenyl]-3-(2-thienyl]-1H-pyrazol-4-yl]methylene]-2-thioxo- (9CI) (CA INDEX NAME)

Searched by Jason M. Nolan, Ph.D.

L7 ANSWER 14 OF 38 CAPLUS COPYRIGHT 2007 ACS ON STN ACCESSION NUMBER: 2004:781003 CAPLUS ON STN 141:288661 DOCUMENT NUMBER: TITLE: INVENTOR(S): 141:298661
Dye-sensitized photoelectric conversion device
Ikeda, Masaaki; Shigaki, Koichiro; Inoue, Teruhisa
Nippon Kayaku Kabushiki Kaisha, Japan
PCT Int. Appl., 75 pp.
CODEN: PIXXD2
Patent
Japanese
1 PATENT ASSIGNEE (S): SOURCE: DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. DATE KIND APPLICATION NO. MO 2004082061

W: AE, AG, AL,
CN, CO, CR,
GE, GH, GH,
LK, LR, LS,
NO, NZ, OK,
TJ, TM, TN,
RW: BW, GH, GH,
BY, KG, KZ,
ES, FI, FR,
KK, TR,
TD, TG
CA 2518825

EP 1628356

R: CH, DE, FR,
CN 1762068

US 2006130249

RITY APPLN. INFO:: Al 20040923 W0 2004-JP3203
AM, AT, AU, AZ, BA, BB, BG, BR, BW,
CU, CZ, DE, DK, DM, DZ, EC, EZ, EG,
HR, HU, ID, ILI, IN, IS, JP, KE, KG,
LT, LU, LV, MA, MD, MG, MK, MN, MM,
PG, PH, PL, PT, RO, RU, SC, SD, SE,
TR, TT, TZ, UA, UG, US, UZ, VC, VN,
KE, LS, MW, MZ, SD, SL, SZ, TZ, TZ, UG,
MD, RU, TJ, TM, AT, BE, BG, CH, CY,
BJ, CF, CG, CI, CM, GA, GN, GQ, GW, 20040311 BZ, CA, CH, FI, GB, GD, KR, KZ, LC, MZ, NA, NI, SK, SI, SY, 2A, ZM, ZW ZW, AM, AZ, DE, DK, EE, RO, SE, SI, MR, NE, SN, A1 20040923 A1 20060222 GB, LI, F1 A 20060419 A1 20060622 CA 2004-2518925 EP 2004-719617 20040311 20040311 CN 2004-80006926 US 2005-548858 JP 2003-70321 20040311 20050909 A 20030314 PRIORITY APPLN. INFO.: A 20030318 JP 2003-73587 W 20040311 WO 2004-JP3203

The title photoelec, conversion device sensitized by organic dye is used preparation of solar cell. Semiconductor micro particles with methine

specified skeleton is used to prepare the photoelec. conversion element which has high conversion efficiency and high applicability. The semiconductor micro particles consist of titania, zinc, or tin. 762269-58-9P

L7 ANSWER 15 OF 38
ACCESSION NUMBER: 2004:430800 CAPLUS
DOCUMENT NUMBER: 140:423667
TITLE: Apreparation of rhodanine derivatives, useful as inhibitors of ubiquitination
INVENTOR(S): Singh, Rajinder; Ramesh, Usha V.; Goff, Dane; Laidig, Guy; Isaakani, Sarkiz D.; Huang, Jianing; Payan,

Donald G.
Rigel Pharmaceuticals, Inc., USA
PCT Int. Appl., 71 pp.
CODEN: PIXXD2 PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: Patent English

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

	PAT	FENT	NO.			KIN		DATE				ICAT					ATE	
	WO	2004	0439					2004	0527								0031	113
		w:	AE,	AG,	AL.	AM,	AT,	AU,	AZ.	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,
			co,	CR,	Cυ,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
			GM,	HR,	HU,	ID,	IL.	IN,	IS,	JP,	KE,	KG,	KP.	KR,	KZ,	LC,	LK,	LR,
			LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NI,	NO,	NZ,	OM,
			PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,	TJ,	TM,	TN,
			TR,	TT,	TZ,	UA,	UG,	US,	υz,	vc,	VN,	YU,	ZA,	ZM,	ZW			
		RW:	BW,	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	AZ,
			BY,	KG,	KZ,	MD.	RU,	TJ,	TM,	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,
			ES,	FI.	FR.	GB,	GR,	HU,	IE,	IT.	LU,	MC,	NL,	PT,	RO,	SE,	SI,	SK,
								CI.										
TG																		
	ΑU	2003	2910	24		A1		2004	0603		AU 2	003-	2910	24		2	0031	113
	EΡ	1597	255			A1		2005	1123		EP 2	003-	7836	09		2	0031	113
		R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
			IE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL,	TR,	BG,	CZ,	EE,	HU,	SK	
	US	2006	2765	20		A1		2006	1207		US 2	005-	5349	19		2	0050	510
PRIO		APP																
											US 2	003-	5149	51P	1	P 2	0031	028

OTHER SOURCE(S): MARPAT 140:423667

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

This invention describes rhodanine derivs. of formula I [wherein: A is [heterojary]: B is C1-6alky] or C2-6alkeny]; X is S, O, etc.; Y is S, O, S[O], or SO2, etc.; R1 = H, NNI2, C1-6alky1, or C1-2alkeny1, etc.; R2 = H, halogan, C1-6alky1, C0-6alky1, theterojary1, or NO2, etc.; R3 = H, C1-6alky1, or C2-6alkeny1; or R3 and B together with the carbon atom to which they are attached form an alkeny1 or a spirocyclic tring, useful as inhibitors of the biochem, pathways of organisms in which ubiquitination is involved. The invention compds. were screened in MDM2 assay (measuring the attachment of ubiquitin to p53) and APC-11/APC-2 ligase assay (auto-ubiquitination). In particular, the compds. and

WO 2003-US36747

W 20031113

ANSWER 14 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

REFERENCE COUNT:

THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

ANSWER 15 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued) compns. are useful for treating cell proliferative diseases such as cancers. For instance, rhodanine deriv. II was prepd. via addn. of Et thioglycolate to benzyl isothiocyanate, intramol. heterocyclization of

obtained carboxylate III, and condensation of furan deriv. IV with the obtained thiazolone V (example 1). 691881-88-6P RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (preparation of rhodamine derivs. and pharmaceutical compns.

(preparation of anomalian delivers of the containing them,
useful as inhibitors of ubiquitination)

RN 691881-88-6 CAPLUS

CL -Phenylalanine, N-[[(52)-5-[[5-(2-chlorophenyl)-2-furanyl]methylene]-4-oxo-2-thioxo-3-thiazolidinyl]acetyl]-, 1,1-dimethylethyl ester (9CI) (CINDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

$$\bigcap_{C1} \bigcap_{S} \bigcap_{N} \bigcap_{N} \bigcap_{Ph} \bigcap_{OBu-t}$$

REFERENCE COUNT: THIS

22 THERE ARE 22 CITED REFERENCES AVAILABLE FOR

RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

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L7 ANSWER 16 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 2004:291950 CAPLUS
DOCUMENT NUMBER: 140:315042
Finl-modulating compounds and methods of use for the treatment of Pinl-associated diseases, including
                                                                      treatment of Fini-associated diseases, including cancer
McKee, Timothy D.; Suto, Robert K.; Tibbitts, Thomas;
Sowadski, Janusz
Pintex Pharmaceuticals, Inc., USA
PCT Int. Appl., 166 pp.
CODEN: PIXXD2
Parent
INVENTOR(S):
PATENT ASSIGNEE(S):
SOURCE:
DOCUMENT TYPE:
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                                                                       Patent
English
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PATENT	NO.	KIND	DATE	APPLICATION NO.	DATE
WO 200	4028535	A1	20040408	WO 2003-US6675	20030303
W:	AE, AG,	AL, AM, AT	AU. AZ.	BA, BB, BG, BR, BY,	BZ, CA, CH, CN,
				DZ, EC, EE, ES, FI,	
	GM. HR.	HU. ID. IL	. IN. IS.	JP, KE, KG, KP, KR,	KZ. LC. LK. LR.
				MK, MN, MW, MX, MZ,	
	PL, PT,	RO, RU, SC	, SD, SE,	SG, SK, SL, TJ, TM,	TN, TR, TT, TZ,
	UA, UG,	UZ, VC, VN	, YU, ZA,	ZM, ZW	
RW	: GH, GM, 1	KE, LS, MW.	, MZ, SD,	SL, SZ, TZ, UG, ZM,	ZW, AM, AZ, BY,
	KG, KZ, I	MD, RU, TJ	, TM, AT,	BE, BG, CH, CY, CZ,	DE, DK, EE, ES,
	FI, FR,	GB, GR, HU	, IE, IT,	LU, MC, NL, PT, RO,	SE, SI, SK, TR,
	BF, BJ,	CF, CG, CI	, CM, GA,	GN, GQ, GW, ML, MR,	NE, SN, TD, TG
AU 200	3225669	A1	20040419	AU 2003-225669	20030303
US 200	4214872	A1	2004102B	US 2003-379408	20030303
EP 155	1396	A1	20050713	EP 2003-798653	20030303
R:	AT, BE,	CH, DE, DK	, ES, FR,	GB, GR, IT, LI, LU,	NL, SE, MC, PT,
	IE, SI,	LT, LV, FI	, RO, MK,	CY, AL, TR, BG, CZ,	EE, HU, SK
PRIORITY AP	PLN. INFO.	:		US 2002-414077P	P 20020926

WO 2003-US6675 W 20030303

OTHER SOURCE(s): MARPAT 140:315042

AB The invention is directed to modulators, e.g., inhibitors, of Pin1 and Pin1-related proteins and the use of such modulators for treatment of associated states, e.g., for the treatment of cancer. Synthetic methods are
included.

IT 312601-58-4 676647-88-4 676654-51-6
RI: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(Pin1-modulating compds. for treatment of Pin1-associated diseases, including cancer)
RN 312601-58-4 CAPLUS
CN 3-Thiazolidinepropanoic acid,
4-oxo-5-[[1-phenyl-3-(2-thienyl)-1H-pyrazol-4-yl]methylene]-2-thioxo- (9CI) (CA INDEX NAME)

ANSWER 16 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

L7 ANSWER 16 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN HO2C - cн₂- сн₂ 676647-88-4 CAPLUS
3-Thiazolidinebutanoic acid, 4-oxo-5-([2,2':5',2''-terthiophen]-5ylmethylene)-2-thioxo- (9CI) (CA INDEX NAME)

RN 676654-51-6 CAPLUS
CN 3-Thiazolidinebutanoic acid,
5-{(3,4-diphenyl-2-furanyl)methylene}-4-oxo-2thioxo- {9CI} (CA INDEX NAME)

(сн2) 3 - со2н

REFERENCE COUNT: THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

L7 ANSWER 17 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
111:136869
Interactions between penicillin-binding proteins (PBPs) and two novel classes of PBP inhibitors, arylalkylidene rhodanines and arylalkylidene iminothiazolidin-4-ones
2ervosen, Astrid; Lu, Wei-Ping; Chen, Zhouliang; White, Ronald E.; Demuth, Thomas P., Jr.; Frere, Jean-Marie
CORPORATE SOURCE: Centre for Protein Engineering, University of Liege, Liege, B-4000, Belg.
SOURCE: Antimicrobial Agents and Chemotherapy (2004), 48(3), 951-969
PUBLISHER: DOCUMENT TYPE: Journal

PUBLISHER: DOCUMENT TYPE: LANGUAGE:

Journal English

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Several non- β -lactam compds. were active against various gram-pos. and gram-neg. bacterial strains. The MIGs of arylalkylidene rhodanines, e.g. I and II, and arylalkylidene iminorhiazolidin-d-ones, e.g. III and IV, were lower than those of ampicillin and cefotaxime for methicillin-resistant Staphylococcus aureus MI339 and

vancomycin-resistant Enterococcus faecium EF12. Several compds. were found to inhibit the

wall synthesis of S. aureus and the last two steps of peptidoglycan biosynthesis catalyzed by ether-treated cells of Escherichia coli or cell wall membrane prepns. of Bacillus megaterium. The effects of the arylalkylidene rhodanines and arylalkylidene inhinthiazolidin-4-one derivs. on E. coli PBP 3 and PBP 5, Streptococcus pneumoniae PBP 2xS (PBP 2x from a penicillin-sensitive strain) and PBP 2xR (PBP 2x from a penicillin-sensitive strain) and PBP 2xR (PBP 2x from a penicillin-resistant strain), low-affinity PBP 2a of S. aureus, and the Actinomadura sp. strain R39 and Streptomyces sp. strain R30 Dp-peptidases were studied. Some of the compds. exhibited inhibitory activities in the 10 to 100 µM concentration range. The inhibition of PBP 2xS by several

them appeared to be noncompetitive. The dissociation constant for the

best inhibitor (Ki = 10 μ M) was not influenced by the presence of the

724784-13-8 724784-14-9 724784-15-0 724784-20-7 724784-20-7 724784-28-5 724784-29-6

724784-20-7 724784-28-5 724784-29-6
RL: BSU (Blological study, unclassified); PAC (Pharmacological activity);
BIOL (Blological study)
(interactions between penicillin-binding proteins and
penicillin-binding protein inhibitors arylalkylidene rhodanines and
arylalkylidene iminothiazolidinones)
724784-13-8 CAPLUS
3-Thiazolidineacetic acid, 5-[[5-(3-chlorophenyl)-2-furanyl]methylene]-4oxo-α-(phenylmethyl)-2-thioxo- (9CI) (CA.INDEX NAME)

(Continued)

(Continued)

ANSWER 17 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

724784-14-9 CAPLUS 3-Thiazolidineacetic acid, $4-oxo-\alpha-(phenylmethyl)-2-thioxo-5-[{5-{3-(trifluoromethyl)phenyl}-2-furanyl]methylene]- {9CI} (CA INDEX NAME)$

724784-15-0 CAPLUS 3-Thiazolidineacetic acid, 4-oxo-5-[[5-(4-phenoxyphenyl)-2-furanyl]methylene]- α -(phenylmethyl)-2-thioxo-(9CI) (CA INDEX NAME)

RN 724784-20-7 CAPLUS
CN Benzoic acid,
2-chloro-5-[5-[2-[[3-[(1,1-dimethylethoxy)carbonyl]phenyl]i
mino]-4-oxo-3-(phenylmethyl)-5-thiazolidinylidene]methyl]-2-furanyl][9CI] (CA INDEX NAME)

L7 ANSWER 17 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

724784-28-5 CAPLUS L-Phenylalanine, N-[5-[[5-(3-chlorophenyl)-2-furanyl]methylene]-4-oxo-3-phenyl-2-thiazolidinylidene]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.
Double bond geometry unknown.

724784-29-6 CAPLUS
3-Thiazolidineacetic acid, 5-[[5-(3-chlorophenyl)-2-furanyl]methylene]a-[(4-hydroxyphenyl)methyl]-4-oxo-2-(phenylimino)- (9CI) (CA INDEX NAME)

ANSWER 17 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

PAGE 2-A

REFERENCE COUNT: THIS

FORMAT

THERE ARE 31 CITED REFERENCES AVAILABLE FOR RECORD. ALL CITATIONS AVAILABLE IN THE RE

L7 ANSWER 18 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2003:948066 CAPLUS

DOCUMENT NUMBER:

TITLE:

INVENTOR (S):

140:22035
Photoelectric converters using dyes with good conversion efficiency
Horiuchi, Tamotsu: Miura, Hidetoshi
Mitsubishi Paper Mills, Ltd., Japan
Jpn. Kokai Tokkyo Koho, 11 pp.
CODEN: JKXXAF
Patent
Japanese 1 PATENT ASSIGNEE (S): SOURCE;

DOCUMENT TYPE: LANGUAGE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE JP 2003346925 PRIORITY APPLN. INFO.: JP 2002-150014 JP 2002-150014 20031205 20020524

OTHER SOURCE(S): MARPAT 140:22035

L7 ANSWER 18 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued) L7 ANSWER 19 OF 38 CAPLUS COPYRIGHT 2007 ACS ON STN ACCESSION NUMBER: 2003:904208 CAPLUS COPYRIGHT 2007 ACS ON STN 2003:904208 CAPLUS CAPLUS DOCUMENT NUMBER: TITLE: 141:33248
Development of a screening assay to measure the loss of antibacterial activity in the presence of its use in optimizing compound structure Roychoudhury, Siddhartha: Brill, Jessica L.; Lu, Wei-Ping; White, Ronald E.; Chen, Zhuoliang; Demuth, Thomas P., Jr.
Clinical Affairs, Ortho-McNeil Pharmaceutical Inc., Hamilton, OH, 45011, USA
Journal of Biomolecular Screening (2003), 8(5), 555-558 proteins: AUTHOR (S): CORPORATE SOURCE: SOURCE . JJJ-JSB CODEN: JBISF3; ISSN: 1087-0571 Sage Publications Journal PUBLISHER: Sage Publications
DOCUMENT TYPE: Journal
LANGUAGE: English
AN assay quantifying the loss of antibacterial potency of compds.,
originally identified via target-based screening, in the presence of
increasing albumin concentration was developed and used as a technique
to measure to measure potential association of compds, with proteins unrelated to their mol. target.

Min. inhibitory concns. (MICs) of test compds, were measured against Staphylococcus aureus strain ATCC 6538 in the presence of 0-12 µM bovine serum albumin (BSA). The linear regression coefficient r2 for the correlation between MIC and BSA concentration was ≥0.9 for 49 and >0.5 62 out of a total of 69 compds. tested. The slope of these correlations varied widely from <1 to 99, suggesting that the loss of potency due to a given concentration of BSA could vary from compound to compound due to variation in the apparent stoichiometry for protein-ligand association Follow-up expts.

using addnl. proteins and a fatty acid, oleic acid, showed that this compound:BSA association was not protein specific, but was likely driven hydrophobicity. The method described in this report can be used to optimize compound design and minimize this undesirable effect. 701979-51-3, FOE 1744694 RE. BSU [Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (development of a screening assay to measure the loss of antibacterial activity in presence of proteins) 701979-51-3 CAPLUS 3-Thiazolidineacetic acid, 5-[[5-(4-chlorophenyl)-2-furanyl]methylene]-a-(hydroxyphenylmethyl)-4-oxo-2-thioxo-, monosodium salt (9CI) (CA INDEX NAME)

ANSWER 19 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

• Na

REFERENCE COUNT: FORMAT

THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

L7 ANSWER 20 OF 38 CAPLUS COPYRIGHT 2007 ACS ON STN

ACCESSION NUMBER:
DOCUMENT NUMBER:
139:307755
Freparation of thiazolidinecarboxamides as prostaglandin FZa receptor modulators
Page, Patrick; Orand-Lebrun, Catherine; Quattropani, Anna; Pomel, Vincent; Schwarz, Matthias; Hamelin, Estelle; Thomas, Russell J.

PATENT ASSIGNEE(S):
Applied Research Systems Ars Holding N.V., Neth. Antilles
SOURCE:
DOCUMENT TYPE:
LANGUAGE:
PAMELY ACC. NUM. COUNT:
FAMILY ACC. NUM. COUNT:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION: DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

WO 2003082278 A1 20031009 WO 2003-EP50083 20030327

W: AZ, AG, AL, AM, AT AU, AZ, BB, BG, BR, BY, BZ, CA, CH, CN, CC, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, BG, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JF, KE, KG, KF, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, ND, MG, MK, MN, MW, MK, MX, NO, NZ, OM, PH, PL, FT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, AZ, AM, ZW

RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AW, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, CA, GW, MC, MR, NE, SN, TD, TG

CA 2477265 A1 20031013 AU 2003-240757 20030327

EP 1487442 A1 20031013 AU 2003-240757 20030327

EP 1487442 A1 20041222 EP 2003-730168

E: AT, BE, CR, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK

BR 2003008748 A 20050111 BR 2003-87486 20030327

JP 2005531524 T 20051020 JP 2003-87186 20030327

JP 2005531524 T 20051020 JP 2003-579816 20030327

JP 2005531524 T 20051020 JP 2003-579816 20030327

JP 200551505 A1 20050929 US 2005-508014 20050512

PRIORITY APPIN. INFO:: EP 2002-100314 A 20020328 PATENT NO. WO 2003-EP50083 W 20030327

OTHER SOURCE(S):

MARPAT 139:307755

ANSWER 20 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

$$(\mathbb{R}^4)_{\,\mathbf{n}} \xrightarrow{\begin{array}{c} \mathbf{S} \\ \mathbf{H} \\ \mathbf{I} \\ \mathbf{I}$$

Title compds. I [wherein G = alkyl(hetero)aryl, alkyl(hetero)cycloalkyl, (hetero)aryl, or (hetero)cycloalkyl which may be fused with cycloalkyl or (hetero)aryl groups; Rl = (hetero)aryl or (heterocycloalkyl which may be fused with (hetero)cycloalkyl or (hetero)aryl groups; R2 = H, (alkyl)carboxy, (alkyl)acyl, (alkyl)alkoxycarbonyl, (alkyl)aminocarbonyl, alkylacyloxy, alkylacylamino, alkylusidido, alkylauifonyloxy, alkylsulfinyl, alkylsulfonyl(maino, alkylsulfonyloxy, alkylsulfinyl, alkylsulfinyl, (hetero)cycloalkyl, alkyl, (hetero)aryl, alkyl, alkenyl, alkyl, alkenyl, alkyl, alkoxy, alkyl, alkenyl, alkyl; or CR2G = cycloalkyl, alkenyl (hetero)aryl, or alkynyl; or CR2G = cycloalkyl; R4 = alkyl, alkenyl, or alkynyl; n = 0-2; geometrical isomers, optically active forms, and pharmaceutically acceptable salts and pharmaceutically active derivs. thereof) were prepared as prostaglandin F2a (PGF2G) receptor modulators. For example, conversion of [1,1'-biphenyl]-4-sulfonic acid

the acid chloride with thionyl chloride, followed by coupling with $N-\{(1S)-3-hydroxy-1-phenylpropyl]-1,3-thiazolidine-2-carboxamide-HCl in the presence of TEA in DCM and chromatog. separation of the$

tereomers
gave (2S)-II and (2R)-II in an overall yield of 58%. (2S)-II exhibited
binding affinity for the human PGF2α receptor with Ki of 0.065 μM
and inhibited inositol triphosphate synthesis and Ca2+ mobilization in
HEK/EBNA cells expressing the human prostaglandin PGF2α receptor
with ICSO values of 0.185 μM and 0.048, resp. PGF2α- or
fluprostenol-induced uterine contractions were reduced by 26% in
non-pregnant rats 40 min after i.v. administration of (2S)-II at a
cumulative dose of 30 mg/kg, and spontaneous uterine contractions were
suppressed by >50% in late-term pregnant rats upon i.v. administration of
(2S)-II over 10 min at a cumulative dose of 30 mg/kg. Thus, I and their
pharmaceutical compns. are useful for the treatment and/or prophylaxis of
preterm labor, premature birth, dysmenorrhea, and for stopping labor r

r to cesarean delivery. 612533-41-2P RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES

ees) (PGF2α ligand; preparation of thiazolidinecarboxamides as prostaglandin F2α receptor modulators for treatment of preterm

ANSWER 20 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued) labor, premature birth, dysmenorrhea, and for stopping labor) 612533-41-2 CAPLUS L-Phenylalanine, N-[[3-{[5-(2-pyridinyl)-2-thienyl]sulfonyl]-2-thiazolidinyl]carbonyl]- (9CI) (CA INDEX NAME)

L7 ANSWER 21 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued) solar cell.

IT 609848-66-0P
RL: PRP (Properties); SPN (Synthetic preparation); TEM (Technical or engineered material use); PREP (Preparation); USES (Uses)
(preparation of thienylpyrrole dye for photoelec. element)
RN 609848-66-0 CAPLUS
CN 3-Thiazolidineacetic acid,
5-[[1-[4-(diethylamtho)phenyl]-2,5-di-2-thienyl1H-pyrrol-3-yl|methylene|-4-oxo-2-thioxo- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

3

REFERENCE COUNT:

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

L7 ANSWER 21 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 2003:773825 CAPLUS
DOCUMENT NUMBER: 139:310012
Photoelectric element with high conversion efficiency from thienylpyrrole dye-sensitized oxide semiconductor microparticle Ikeda, Masaaki: Shigaki, Koichiro; Inoue, Teruhisa Nippon Kayaku Co., Ltd., Japan Jpn. Kokai Tokkyo Koho, 28 pp. CODEN: JKXXAF INVENTOR(S): PATENT ASSIGNEE(S): SOURCE: DOCUMENT TYPE: Patent LANGUAGE: Japanese FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. DATE JP 2003-7360 JP 2002-7062 JP 2003282165 PRIORITY APPLN. INFO.: 20031003 20030115 A 20020116 MARPAT 139:310012 OTHER SOURCE(S):

AB The photoelec. element comprises an oxide semiconductor microparticle thin film sensitized by a dye I (R, Z = H, alkyl, aromatic hydrocarbon

fue, heterocyclyl; Y1, Y2 = substituent may form ring; and m1, m2 = integer 0-3). Alternatively, the dye is represented by II (A1, A2 = aromatic hydrocarbon residue, heterocyclyl, etc.; X1 = aromatic hydrocarbon

due, heterocyclyl, cyano, etc.; and $l=integer\ l-3$). The oxide semiconductor microparticle may be made from TiO2. The photoelec. element is used for

Searched by Jason M. Nolan, Ph.D.

L7 ANSWER 22 OF 38 CAPLUS COPYRIGHT 2007 ACS ON STN ACCESSION NUMBER: 2003:719505 CAPLUS DOCUMENT NUMBER: 139:240341 Finl peptidyl prolyl isomerase-heterocyclic

Pin1 peptidyl prolyl isomerase-modulating

compounds and methods of use for the treatment of cancer and other Pinl-associated conditions McKee, Timothy D.; Suto, Robert K.
Pintex Pharmaceuticals, Inc., USA PCT Int. Appl., 79 pp.
CODEN: PIXED2
Patent
English

INVENTOR(S):
PATENT ASSIGNEE(S):
SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: English

P	TENT	NO.			KIN	D	DATE			APPL	ICAT	ION	NO.		D	ATE	
						-									_		
W	200	30745	50		A2		2003	0912		WO 2	003-	US 63	94		2	0030	303
wo	200	30745	50		A3		2003	1204									
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		co,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
		GM,	HR,	ΗU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	ΚZ,	LC,	LK,	LR,
		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	OM,	PH,
		PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	sĸ,	SL,	TJ,	TM,	TN,	TR,	TT,	TZ,
		UA,	UG,	UZ,	VÇ,	VN,	YU,	ZA,	ZM,	2W							
	RW	: GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	AZ,	BY,
		KG,	KZ,	MD,	RU,	ТJ,	TM,	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,
		FI,	FR,	GB,	GR,	HU,	IE,	IT,	LU,	MC,	NL,	PT,	RO,	SE,	SI,	SK,	TR,
		BF,	BJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG
A	200	32136	73		Al		2003	0916		AU 2	003-	2136	73		2	0030	303
U	200	41763	72		A1		2004	0909		US 2	003-	3793	77		2	0030	303
PRIORI	Y AP	PLN.	INFO	.: ·						US 2	002-	3612	06P		P 2	0020	301
										WO 2	003-	US 63	94	,	w 2	0030	303

OTHER SOURCE(S):

MARPAT 139:240341

ANSWER 22 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued) thienyl)-1H-pyrazol-4-yl]methylene}-4-oxo-2-thioxo- (9CI) (CA INDEX

ANSWER 22 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

AB The invention discloses modulators, e.g. inhibitors, of Pin1 and Pin1 related proteins, and the use of such modulators for treatment of Pin1 associated states, e.g., for the treatment of cancer. Modulators of the invention include I [dashed lines = single or double bond; n = 0-2; G1 = CH2, CH, N; G2, G3 = H, alkyl, aryl, O, etc.; R1-R7 = H, acyl, (un)substituted alkyl, etc.]. Determination of Pin1 overexpression in a Variety of tumor types is also presented.

1T 596807-21-5 596807-21-5D, derivs.
RL: PAC (Pharmacological activity); TRU (Therapeutic use); BIOL (Biological study); USES (Uses) (Pin1 peptially prolyl isomerase-modulating heterocyclic compds. for treatment of cancer and other Pin1-associated conditions)

RN 596807-21-5 CAPULS
CN 3-Thiazolidinepropanoic acid, 5-[[5-hydroxy-1-(4-methylphenyl)-3-(2-thienyl)-1H-pyrazol-4-yl]methylene]-4-oxo-2-thioxo- (9CI) (CA INDEX NAME)

596807-21-5 CAPLUS
3-Thiazolidinepropanoic acid, 5-[{5-hydroxy-1-(4-methylphenyl)-3-(2-

L7 ANSWER 23 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
139:6866
Preparation of 5-(benzylidene)rhodanines and analogs as antidiabetics and antitumor agents
Pfahl, Magnus; Tachdjian, Catherine; Spruce, Lyle W.;
Al-Shamma, Hussien A.; Boudjelal, Mohamed; Fanjul, Andrea N.; Wiemann, Torsten R.; Pleynet, David P. M.
PATENT ASSIGNEE(S):
SOURCE:
CODEN: PIXAD2
Patent INFORMATION:

1

CAPLUS COPYRIGHT 2007 ACS on STN
2003:417731 CAPLUS
Preparation of 5-(benzylidene)rhodanines and analogs as antidiabetics and antitumor agents
An Hussien A.; Boudjelal, Mohamed; Fanjul, Andrea N.; Wiemann, Torsten R.; Pleynet, David P. M.
Maxia Pharmaceuticals, Inc., USA
PCT Int. Appl., 118 pp.
CODEN: PIXAD2
Patent INFORMATION:
1

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

	PENT I				KIN	D	DATE			APPL	ICAT	ION	NO.		D.	ATE	
	2003						2003	0530		WO 2	002-	US36	583			0021	
	W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN
		co,	CR,	CU,	cz,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH
		GM,	HR,	ΗU,	ID,	IL,	IN,	ĮS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR
		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	OM,	PH
		PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	TJ,	TM,	TN,	TR,	TT,	TZ
		UA,	UG,	UZ,	VN,	YU,	ZA,	ZM,	ZW								
	RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	AZ,	BY
		KG,	KZ,	MD,	RU,	TJ,	TM,	AT,	BE,	BG,	CH,	CY,	cz.	DE,	DK,	EE,	ES
		FI.	FR.	GB.	GR,	IE,	IT,	LU,	MC.	NL.	PT.	SE,	sK,	TR.	BF,	BJ,	CF
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ΑIJ	2002														2	0021	115
us	2003	1443	29		A1		2003	0731		US 2	002-	2980	24			0021	
	7071				B2		2006	0704							-		
	1456				A1		2004	0915		EP 2	002-	7896	54		2	0021	115
			RE.	CH.				FR,									
								MK,								,	••
JP.	2005		26	,	т,	,	2005	0512	0.,	.TP 2	003-	5456	35	,	,	0021	115
119	2003	2164	32		Δ1		2003	1120		115 2	003-	3843	52		2	0020	306
US	2003 7102	000	-		R2		2006	0905			~~~				-	0000	-
211	2004	N340	n.a		14		2004	0219		119 2	003-	3843	91		,	0030	306
115	7196	108	••		R2		2007	0227		00 L	005	3043			-	0000	300
	2478				B2 A1 A1		2003	0327 0918		CA 2	003-	2478	342		,	0030	207
	2478				A1		2003	0918		CA 2	003-	2478	765		5	0030	307
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								IS,									
		LS										MX,	MZ	NT		NZ.	

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ANSWER 23 OF 38 CAPLUS COPYRIGHT 2007 ACS ON STN (Continued)
TZ, UA, UG, UZ, VC, VN, YU, ZA, ZM, ZW
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,
KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES,
FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR,
BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
AU 2003223233 Al 20030922 AU 2003-223233 2003007
AU 2003223233 A2 20030922
AU 2003222682 A2 20030922
AU 2003225682 A2 20030922
AU 2003225682 A1 2003092
EP 1487446 A2 20041222 EP 2003-719363 200307
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, LT, LI, LU, NL, SE, MG, PT,
          BR 2003008278
CN 1649586
JP 2005525371
JP 2005530705
NO 2004004250
US 2006160796
US 2006241138
PRIORITY APPLN. INFO.:
                                                                                                             US 2001-334794P
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                                                                                                                                                              A3 20030306
                                                                                                             WO 2003-US6784
                                                                                                                                                              W 20030307
                                                                                                             WO 2003-US7240
                                                                                                                                                                   20030307
OTHER SOURCE(S):
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MARPAT 139:6866

ANSWER 23 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued) to controls. In addn., III displayed selective potency against various human cancer cell lines; e.g. at a conen. of 10 μ M, about 80% of breast cancer cells were killed compared to ≤ 50 % of other cell lines studied. Thus, I and II are useful in the treatment of diseases, such

cancer, metabolic disorders, Type 2 Diabetes, dyslipidemia, and/or hypercholesterolemia. 532440-69-0P RE: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES

es; (antidiabetic and/or antitumor agent; preparation

(antidianetic district and the first and the first and the diseases) (benyilidene) rhodanines and analogs for treatment of cancer, diabetes, and other diseases) RN 532440-69-0 CAPPLUS CN 3-Thiazolidineacetic acid, 5-{[5-(6-methoxy[1,1'-bipheny1]-3-y1}-2-furany1]methylene]-4-oxo-2-thioxo-(9CI) (CA INDEX NAME)

REFERENCE COUNT:

FORMAT

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

ANSWER 23 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

Title benzylidene-substituted 2-thioxo-4-oxothiazolidines and analogs I and II [wherein Ar1 = 2-(R7)-4-(R5)-5-(R6)C6H2 optionally substituted

and II (wherein Ar1 = 2-(R7)-4-(R5)-5-(R6)CGH2 optionally substituted with R8; Ar2 = (hetero)aryl; W = S, O, or NR3; X = O or S; R1 = H or (un)substituted organic radical comprising 1-4 C's; R2 = (un)substituted organic radical comprising 1-12 C's; R3 = H or (un)substituted organic radical comprising 1-12 C's; C2R5R6 = 5-7 membered non-aromatic ring optionally comprising 1-2 heteroatoms; R7 and R8 = independently H or (un)substituted alkyl or amino; and pharmaceutically acceptable salts thereof) were prepared as 1 liver X receptor (LXR), peroxisome proliferator-activated receptor 7 (PPARY), protein kinase Akt/PKB (AKT-1/PKBa) inhibitors. For example, esterification of 6-hydroxynaphthoic acid with EtOH (98%), followed by protection with triflic anhydride in CH2C12 gave 6-(trifluoromethanesulfonyloxy)naphthalene-2-carboxylic acid Et ester (100%). Reduction of the ester to the alc. (72%) using DIBAL, conversion to the aldehyde (94%) with PCC, and Suzuki coupling with (3-dimethylamino-5,5,8,8-tetramethyl-5,6,7,8-tetrahydronaphthalene-2-carboxaldehyde (71%).

Coupling of the aldehyde with rhodanine-3-acetic acid in the presence of piperidine and acetic acid in toluene afforded III (33% yield, 99.5% purity). The latter antagonized both LXR and PPRRy activation in vitro in a dose-dependent fashion, reaching inhibition values of about 80%-90% at 10 µM. Oral administration of III to rats maintained on a high cholesterol atherogenic diet resulted in significant redus. In total serum cholesterol and low d. lipoprotein cholesterol levels with accompanying elevations in high d. lipoprotein cholesterol levels arred compared

L7 ANSWER 24 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 2003:143481 CAPLUS
DOCUMENT NUMBER: 138:195809 High sensitive silver halide photographic material containing methine dye with linked chromophores (ADDAMSAME, SUBJEUT, TAKIZAWA, HIFOO PATEMT ASSIGNEE(S): 50URCE: Upi Photo Film Co., Ltd., Japan JODCUMENT TYPE: CODEN: JKXXAF PATEMT INFORMATION: 1400 ADDAMSAME ADDAMSAM

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE JP 2003057777 PRIORITY APPLN. INFO.: 20030226

The material has ≥ 1 photosensitive Ag halide emulsion layer containing the methine dye Dlr2(LlqaD2q1)r1·CIy [Ll = linkage; Dl, D2 = chromophore; ql, rl, r2 = 1-4; [ql + rl + r2] = 3-5; qa = 1-4; CI = charge neutralizing ion; y = the number for charge neutralization]. 499770-01-3P

RL: PNU (Preparation, unclassified); TEM (Technical or engineered

rial
use); PREP (Preparation); USES (Uses)
(novel methine dye with linked chromophores for photog. sensitizer)
499770-01-3 CAPLUS
Benzothiazolium, 3-[2-[2-[2-[{]3-[2-[2-[3-(carboxymethyl)tetrahydro2,4,6-trioxo-5-[[3-(4-sulfobutyl)naphth[2,1-d]oxazol-2(3H)-

ylidene]ethylidene]-1(2H)-pyrimidinyl]ethoxy]ethoxy]ethyl]tetrahydro-2,4,6trioxo-5-[(3-(4-sulfobutyl)naphth[2,1-d]oxazol-2(3H)-ylidene]ethylidene]1(2H)-pyrimidinyl]acetyl]amino]ethoxy]ethoxy]ethyl]-2-[[5-phenyl-3-[3sulfopropyl)-2(3H)-benzothiazolylidene]methyl]-5-(2-thlenyl)-, inner

disodium salt (9CI) (CA INDEX NAME)

ANSWER 24 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

(Continued)

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ANSWER 24 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

(Continued)

PAGE 1-A

PAGE 1-B

● 2 Na

ANSWER 24 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

(Continued)

PAGE 2-B

ΙŤ 499770-03-5

499770-03-5
RE: TEM (Technical or engineered material use); USES (Uses)
(novel methine dye with linked chromophores for photog. sensitizer)
499770-03-5 CAPLUS
Benzothiazolium, 3-[5-[[[3-[2-[{6-[3-butyl-5-[[1-butyl-3(carboxymethyl)hexahydro-2,4,6-trioxo-5-pyrimidinyl]methylene]tetrahydro-

,6-trioxo-1(2H)-pyrimidinyl]-1-oxohexyl|amino|ethyl]amino|-2-oxoethyl]5-[(3-ethyl-5-sulfo-2(3H)-benzoxazolylidene)ethylidene)tetrahydro-2,4,6trioxo-1(2H)-pyrimidinyl)acetyl]amino|pentyl]-2-[(5-phenyl-3-(3-sulfopropyl)-2(3H)-benzothiazolylidene|methyl]-5-(2-thienyl)-, inner

salt, disodium salt (9CI) (CA INDEX NAME)

L7 ANSWER 25 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 2003:69123 CAPLUS
DCUMENT NUMBER: 138:144993
New methine dye and silver halide photographic material containing the same KODBAysahi, Suguru: Takizawa, Hiroo FOMERIT ASSIGNEE(S): 50URCE: 701 Photo Film Co., Ltd., Japan Jpn. Kokai Tokkyo Koho, 36 pp.
CODEN: JKXXAF
DCUMENT TYPE: LANGUAGE: PANILY ACC. NUM. COUNT: 1

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. APPLICATION NO. KIND DATE DATE JP 2003029366 PRIORITY APPLN. INFO.: 20030129 JP 2001-218231 JP 2001-218231

OTHER SOURCE(S): MARPAT 138:144993

The invention relates to a silver halide photog, material comprised of a support at least one Ag halide photog, emulsion layer, wherein the Ag halide photog, emulsion layer contains the new methine dye represented by pyel-(LI-(Dye2)ml)m2·(CII)yl (LI = connecting group; ml = 1-5; m2 = 1-5; CI = counter ion; yl = number to neutralize; Dyel = first chromophore represented by I; Dye2 = second chromophore; Xl, X2 = 0, S, NR6, CR7R8; R6-8 = H, alkyl, alkenyl, aryl, heterocyclyl; R1, R2 = H, alkyl, alkenyl, aryl, heterocyclyl; M1-3 = methine; nl = 0-3; n2, n3 = 0-4; vl, V2 = substituent; L1 connects to Dyel via R1, R2, V1, or V2]. The photog. material shows improved sensitivity.
492453-13-1 492453-14-2

R1: MOA (Modifier or additive use); USES (Uses)
(new methine dye as spectral sensitizer in silver halide photog. material to improve sensitivity)
492433-13-1 CAPLUS
Benzothiazolium, 3-[6-[[6-[2-[(3-ethyl-6-sulfobenzothiazolium-2-

yl)methylene]-6-sulfo-3(2H)-benzothiazolyl]-1-oxohexyl]oxy]hexyl]-2-[[3-(3-

sulfopropyl)-5-(2-thienyl)-2(3H)-benzothiazolylidene]methyl}-5-(2-thienyl), bis(inner salt), sodium salt (9CI) (CA INDEX NAME)

ANSWER 25 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

PAGE 2-A

492453-14-2 CAPLUS
Benzothiazolium, 3-[2-[2-[2-[[6-[2-[(3-ethyl-6-sulfobenzothiazolium-2-y]]methylene]-6-sulfo-3[2H]-benzothiazolyl]-1exohexyl]oxylethoxylethoxylethyl]-2-[[3-(3-sulfopropyl)-5-(2-thienyl)2[3H]-benzothiazolylidene|methyl]-5-(2-thienyl)-, bis(inner salt), sodium
salt (9CI) (CA INDEX NAME)

L7 ANSWER 25 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

PAGE 1-A

PAGE 1-B

L7 ANSWER 26 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:
DOCUMENT NUMBER: 137:325416

FITTLE: 137:325416

Preparation of substituted imidazoles/oxazoles/thiazoles as large conductance calcium-activated K channel openers

Hongu, Mitsuya; Hosaka, Thoshihiro; Kashiwagi, Toshihiko; Kono, Rikako; Kobayashi, Hiroyuki

Tanabe Seiyaku Co., Ltd., Japan
PCT Int. Appl., 302 pp.
COOEN: PIXXD2

DOCUMENT TYPE: Patent
LANGUAGE: English

DOCUMENT TYPE: LANGUAGE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

		ENT															DATE	
	WO		0831	11		A2		2002	1024	1							20020	
	WO										D7	CB	CN	-	CD	CII	, cz,	DM
		w:	AE,	AG,	AL,	AU,	BA,	вв,	BG,	BK,	BZ,	CA,	CN,	٠٠,	UK,		,,	UM,
			DZ,	EC,	EE,	GD,	GE,	HR,	HU,	In,	IL,	IN,	ıs,	UP,	AK,	TIC.	, LK,	DK,
											NZ,	OM,	PH,	РЬ,	RO,	36	, SI,	SK,
								VN,										D1/
		RW:															, AZ,	
			KG,	KZ,	MD,	RU,	TJ,	TM,	AT,	BE,	CH,	CY,	DE,	DK,	ES,	CI	, FR,	CB,
													Вυ,	CF,	CG,	CI	, см,	GA,
			GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG						20020	415
	CA	2444	596			Al		2002	1024		CA 2	002-	2444	296			20020	415
	AU	2002	2463	97		Al		2002	1028		AU 2	002-	2463	91			20020 20020 20020 20020 20020	413
	HU	2003	0382	9		A2		2004	0301		HU Z	003-	3829				20020	415
	CN	1503	786			A		2004	0609		CN 2	002-	8083	70			20020	415
	EΡ	1432	690			A2		2004	0630		EP 2	002-	7145	77			20020	415
																	, мс,	
			IE,	sı,	LT,	LV,	FI,	RO,	MK,	CY,	AL,	TR						
	ВŔ	2002	0089	56		A		2004	0713		BR 2	002-	8956				20020	415
	JР	2004	5315	22		T		2004	1014		JP 2	002-	5009	15			20020	415
	ΝZ	5290	43	_		A.		2006	1130		NZ Z	002-	5290	43			20020 20020 20020 20040 20050 20010	415
	US	2004	1275	27		A1		2004	0701		US 2	004-	4748	50			20040	210
	AU	2005	2027	51		Al		2005	0714		AU 2	005-	2027	51		_	20050	623
PRIO	RIT	(APP	LN.	INFO	.:						JP 2	001-	1164	36		A	20010	416
																	20010	
											AU 2	002-	2463	97		A3	20020	415
	•										WO 2	002-	JP37	23		w	20020	415

OTHER SOURCE(S): MARPAT 137:325416 L7 ANSWER 26 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

The title compds. [I; X = NR4, O, S; R1, R2 = H, halo, CO2H, etc.; R3 = aryl, heterocyclyl, alkyl; R4 = H, alkyl), useful in the prophylaxis and/or treatment for pollakiuria or urinary incontinence, were prepared Thus, reacting 5-ethyl-2-iodo-4-(3-pyridyl)imidazole with 3-(hydroxymethyl)thiophene-2-boric acid in the presence of Pd(PPh3)4 and aqueous 2M Na2CO3 in dimethoxyethane afforded I.2HCl [X = NH; R1 = Et;

3-pyridyl; R3 = 3-(hydroxymethyl)thien-2-yl] which showed 100% inhibition time of 10-20 min in test on the rhythmic bladder contractions induced by substance P in anesthetized rats.
473691-96-2P 473691-99-5P
RL: PAC (Pharmacological activity); SFN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES

(Uses)
(preparation of imidazoles/oxazoles/thiazoles as large conductance calcium-activated K channel openers)
473691-96-2 CAPLUS
4-Oxazoleacetic acid, 2-[6-(3-thiazolidinyl)-3-pyridinyl]-5-(3-thienyl)-, ethyl ester (9CI) (CA INDEX NAME)

473691-99-5 CAPLUS
4-Oxazoleacetic acid, 2-(6-(3-thiazolidinyl)-3-pyridinyl)-5-(3-thienyl)-, sodium salt (9CI) (CA INDEX NAME)

ANSWER 26 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

• Na

ANSWER 27 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN
JP 2004520839 T 20040715 JP 2002-570761
NZ 528644 A 20050527 NZ 2002-528644
US 2003232816 A1 20031218 US 2002-238282
US 6794404 B2 20040921
US 2004101026 A1 2004610 US 2004-471164
US 2004224955 A1 2004111 US 2004-451710
US 20042161048 (Continued) 20020306 20020306 20020910 US 2004-471164 US 2004-851710 US 2000-216084P US 2004224955 PRIORITY APPLN. INFO.: 20040521 P 20000706 US 2001-274374P P 20010308 US 2001-281343P P 20010405 US 2001-898297 A3 20010703 WO 2001-CA989 W 20010704 US 2001-995099 A3 20011127 WO 2002-CA323 W 20020306 US 2002-238282 A1 20020910

MARPAT 136:118447

(CH2) nCYZ

OTHER SOURCE(S):

Title compds. [I; X = CH, N; Y = O, S; Z = OH, NH2; NMeR3, NHR3, OR3, 5-6 membered (substituted) heterocyclyi A = N, COR7, CR5, R5 = H, halo, alkyl: Y and A are not both N; R6 = H, halo, alkyl, OR7;

= H , alkyl; R1 = (substituted) hetero(bi)cyclyl, Ph, phenylalkyl, alkenyl, phenylalkenyl, cycloalkyl, alkyl, CF3; R2 = (substituted) alkyl, cycloalkyl, cycloalkyl, bicycloalkyl, adamantyl, Ph, pyridyl; R3 =

H,

alkyl, cycloalkyl, cycloakylakyl, arylakyl, arylakyl, alkenyl,
cycloalkylakenyl, arylalkenyl, dialkylamino, heterocyclyl, etc.; n = 0,
1], were prepared Thus, Me 3-amino-4-cyclohexylaminobenzoate
(preparation
given), 2-pyridinecarboxaldehyde, and Oxone were stirred in DMF to give
801 % 1 -cyclohexyl-2-pyridin-2-yl-1H-benzimidazole-5-carboxylate, which
was saponified with aqueous NaOH in MeOH to give 91%
1-cyclohexyl-2-pyridin-2-yl1H-benzimidazole-5-carboxylic acid. The latter inhibited hepatitis C
virus RNA dependent polymerase (NS5B) with IC50 = 1-5 µM.
390811-18-4P
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
(Uses)

(preparation of benzimidazolecarboxylates and related compds. as viral polymerase inhibitors)
390811-18-4 CAPLUS

L7 ANSWER 27 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2002:51438 CAPLUS COPYRIGHT 2007 ACS ON STN 136:118447

DOCUMENT NUMBER: TITLE:

136:1844/
Preparation of benzimidazolecarboxylates and related compounds as viral polymerase inhibitors
Beaulieu, Pierre Louis; Fazal, Gulrez; Gillard, Kukolj, George; Austel, Volkhard Boehringer Ingelheim (Canada) Ltd., Can. PCT Int. Appl. 322 pp. CODEN: PIXXD2 Patent English INVENTOR (S):

PATENT ASSIGNEE (S):

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PAT	ENT:	ΝО.			KIN	D	DATE			APPL	ICAT	ION	NO.		Di	ATE	
															-		
WO	2002						2002									0010	
	W:	ΑE,	AG,	AL,	AM,	ΑT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,
		CR,	CU,	cz,	DE,	DK,	DM,	DZ,	EE,	ES,	FI,	GB,	GD,	GΕ,	GH,	GM,	HR,
		ΗU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	ΚZ,	LC,	LK,	LR,	LS,	LT.
							MK,										
		SD,	SE,	SG,	SI,	SK,	SL,	TJ,	TM,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	٧N
		YU,	ZA,	ZW													
	RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ.	UG,	ZW,	AT,	BE,	CH,	CY.
		DE.	DK.	ES.	FI.	FR.	GB,	GR.	IE.	IT.	LU.	MC.	NL.	PT.	SE.	TR.	BF
		BJ,	CF,	CG,	CI,	CM,	GA,	GN,	GW.	ML,	MR,	NE,	SN,	TD,	TG		
US	2002	0654	18		A1		2002	0530		US 2	001-	8982	97		21	0010	703
US	6448	281			A1 B2		2002	0910									
CA	2412	718			A1		2002	0117		CA 2	001-	2412	718		2	0010	704
ΕP	1301	487			A2		2003	0416		EP 2	001-	9512	74		2	0010	704
EΡ	1301	487			B1		2006	1122									
	R:	AT.	BE.	CH.	DE.	DK.	ES,	FR.	GB.	GR,	IT.	LI.	LU.	NL.	SE.	MC.	PT
		TE.	SI.	LT.	I.V.	FT.	RO.	MK.	CY.	AI.	TR						
JP	2004 3460 6479 2439 2002	5027	61		T		2004	0129	٠.	JP 2	002-	5092	92		2	0010	704
ΑT	3460	49			T		2006	1215		AT 2	001-	9512	74		2	0010	704
US	6479	508			В1		2002	1112		US 2	001-	9950	99		2	0011	127
CA	2439	176			A1		2002	0912		CA 2	002-	2439	176		2	0020	306
WO	2002	0707	39		A2		2002	0912		WO 2	002-	CA32	3		2	0020	306
WO	2002	0707	39		A3		2003	0530							_		
	W:	AE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA.	BB,	BG.	BR.	BY.	BZ.	CA.	CH.	CN
							DK,										
		GM,	HR,	HΨ,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR
		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	OM,	PH
		PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	sĸ,	SL,	TJ,	TM,	TN,	TR,	TT,	TZ
							Yυ,										
	RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	ŪĠ,	ZM,	ZW,	AM,	AZ,	BY
		KG,	KZ,	MD,	RU,	TJ,	TM,	AT,	BE,	CH,	CY,	DE,	DK,	ES,	FI,	FR,	GB
		GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	TR,	BF,	BJ,	CF,	CG,	CI.	CM,	GA
		GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG							
ΑU	2002	2445	66		A1		2002	0919		AU 2	002-	2445	66		2	0020	306
ΕP	1370	682			A2		2003	1217		EP 2	002-	7126	81		2	0020	306
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC.	PT
							RO,					-,			,	,	
HTI	2004						2004					39			2	0020	306

ANSWER 27 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued) L-Phenylalanine, N-[[1-cyclohexyl-2-(3-furanyl)-1H-benzimidazol-5-yl]carbonyl]-4-[(4-oxo-2-thioxo-5-thiazolidinylidene)methyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry unknown.

L7 ANSWER 28 OF 38 CAPLUS COPYRIGHT 2007 ACS ON STN ACCESSION NUMBER: 1999:265975 CAPLUS DOCUMENT NUMBER: 130:289169

DOCUMENT NUMBER: TITLE:

130:289169
Color photographic silver halide material with improved sensitivity and reduced yellow fog Weimann, Ralf; Misfeldt, Michael; Geiger, Markus Agfa-Gevaert Ag, Germany Ger. Offen., 30 pp.
CODEN: GWXXEX INVENTOR(S):

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE :

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 19745886	A1	19990422	DE 1997-19745886	19971017
US 6010838	A	20000104	US 1998-167476	19981007
JP 11212206	A	19990806	JP 1998-307807	19981015
PRIORITY APPLN. INFO.:			DE 1997-19745886 A	19971017

OTHER SOURCE(S):

MARPAT 130:289169

In the color photog, material comprising a support, at least 1 red-sensitive Ag halide emulsion layer containing a cyan coupler, at

green-sensitive Ag halide emulsion layer containing a magenta coupler, at least 1 blue-sensitive Ag halide emulsion layer containing a yellow

coupler,
and optional further layers, the Ag halide contains at least 95 mol.%
AgCl, the blue-sensitive layer is spectral sensitized by I (R1 =
2-thienyl, 3-thienyl; R2, R3 = alkyl, sulfoalkyl, carboxyalkyl,
(CH2)noONHSO2-alkyl, (CH2)nsO2NHSO2-alkyl, (CH2)ncONHSO2-alkyl,
(CH2)ncONHCO-alkyl; R4, R5 = H, halo, alkyl, methoxy, aryl, 2-furanyl,
3-furanyl, 2-thienyl, 3-thienyl, 1-pyrrolyl, 2-pyrrolyl, 3-pyrrolyl,

ANSWER 28 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued) 1-indolyl, N-carbazolyl, 2-isoindolyl; R4 joining together with R5 may form benzene or naphthalene ring; Z = O, S; n = 1-6; X = counterion), and contains a yellow coupler II (R1-3 = alkyl; R2 joining together with R3 may form 3- to 6-membered ring; R4 = alkyl, cycloalkyl, aryl; R5 = halo, alkyl, alkoxy, aryloxy, alkoxycarbonyl, alkylsulfonyl, alkylsurfonyl, arylcarbamoyl, arylcarbamoyl, alkylsulfonamido, arylsulfonamido; m = O-3; Z1 = O, NR6; Z2 = NR7, CR8R9; R6-9 = H, substituent). By combining the yellow coupler and blue sensitizer, the color photog, material shows improved blue-sensitivity

and reduced yellow fog. 222534-78-3

τT

222534-76-3

RL: MCA (Modifier or additive use); USES (Uses)
(blue-sensitive layer of color photog. silver halide material spectrally sensitized with)
222534-78-3 CAPLUS
Benzothiazolium, 3-(2-carboxyethyl)-2-{[3-(3-sulfopropyl)-5-(2-thienyl)-2(3H)-benzothiazolylidene]methyl)-5-(3-thienyl)-, inner salt (9CI) (CA INDEX NAME)

ANSWER 29 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN SSION NUMBER: 1998:799977 CAPLUS MENT NUMBER: 130:38375

ACCESSION NUMBER DOCUMENT NUMBER:

Preparation of 5-furfurylidene-4-thiazolidinones and analogs as vascular endothelial growth factor TITLE:

receptor

....agoniats
Scott, Ian L.; Biediger, Ronald J.; Market, Robert V.
Texas Biotechnology Corporation, USA
PCT Int. Appl., 41 pp.
CODEN: PIXXD2
Patent
English INVENTOR (5): PATENT ASSIGNEE (S):

SOURCE:

DOCUMENT TYPE:

LANGUAGE: English

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PA1	ENT	NO.			KIN	D	DATE			APPL	ICAT	ION	NO.		D	ATE	
						-											
WO	9853	790			A2		1998	1203		WO 1	998-	US 93	66		1:	9980	601
WO	9853	790			A3		1999	0304									
	W:	AL,	AM,	AT,	ΑU,	AZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CU,	CZ,	DE,
		DK,	EE,	ES,	FI,	GB,	GE,	HU,	IL.	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,
		LK,	LR,	LS,	LT,	LU,	LV,	MD,	MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,	PT,
		RO,	RU,	SD,	SE,	SG,	SI,	SK,	TJ,	TM,	TR,	TT,	UA,	UG,	US,	UZ,	VN,
							MD,										
	RW:						SD,				AT.	BE,	CH.	CY.	DE.	DK.	ES,
							IT,										
							NE.						•				
PRIORITY	APP				,	,				US 1	997~	4810	5 P		P 1	9970	530

OTHER SOURCE(S):

MARPAT 130:38375

$$R^1$$
 R^2
 R^3
 Y
 Z
 $N-R$

Title compds. [I: R = TR4: X = O, S, CR5:CR6: R1-R6 = H, cycloalkyl, heterocyclyl, aryl, etc.: R4 = H, (cyclo)alkyl, heterocyclyl, aryl, etc.: T = bond, alkylene, {alkyl}imino, NHCO, etc.: Y = O, S, {alkyl}imino,

CH2;
Z = CH2, CO, CS] were prepared as vascular endothelial growth factor receptor antagonists (no data). Thus, 3-benzyl-4-thiazolidinone was acylated by Me 5-phenyl-2-furoate (preparation each given) and the product
converted in 2 steps to I (R = CH2Ph, R1 = Ph, R2 = R3 = H, X = O, Y = S,

Z = CH2).

17 216772-06-49 216773-13-6P
RL: BAC (Biological activity or effector, except adverse); BSU
(Biological

study, unclassified): SPN (Synthetic preparation): THU (Therapeutic use): BIOL (Biological study): PREF (Preparation): USES (Uses) (preparation of 5-furfurylidene-4-thiazolidinones and analogs as vascular

Searched by Jason M. Nolan, Ph.D.

L7 ANSWER 29 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN endothelial growth factor receptor antagonists)
RN 216772-06-4 CAPLUS (Continued)

Benzenebutanoic acid,
-[{5-(4-chlorophenyl)-2-furanyl]methylene]-4-oxo-2-thioxo-3-thiazolidinyl]- (9CI) (CA INDEX NAME)

216773-13-6 CAPLUS Benzoic acid, 5-[5-[5-[4-[2-(3-cyanophenoxy)ethoxy]pheny1]-2-furanyl]methylene]-4-oxo-2-thioxo-3-thiazolidinyl]-2-[[(2-(2-methoxyethoxy)ethoxy]acetyl]amino]- (9CI) (CA INDEX NAME)

- CH2- O- CH2- CH2- O- CH2- CH2- OM6

L7 ANSWER 30 OF 38 CAPLUS COPYRIGHT 2007 ACS ON STN ACCESSION NUMBER: 1998:298195 CAPLUS DOCUMENT NUMBER: 129:47355 THITLE: Thermally developeble 2009

INVENTOR (S):

129:47355
Thermally developable silver halide photographic material containing cyanine sensitizing dye Inagaki, Yoshio: Tauzuki, Hirohiko Fuji Photo Film Co., Ltd., Japan Jpn. Kokai Tokkyo Koho, 25 pp.
CODEN: JOXXAF PATENT ASSIGNEE (S): SOURCE:

DOCUMENT TYPE:

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 10123663	A	19980515	JP 1996-293332	19961015
JP 3681840	B2	20050810		
US 5998125	A	19991207	US 1997-949694	19971015
PRIORITY APPLN. INFO.:			JP 1996-293332 A	19961015

OTHER SOURCE(S): MARPAT 129:47355
GI For diagram(s), see printed CA Issue.
AB The title materials contain, on ≥1 side of a support,
photosenstive Ag halide grains and mono-, tri-, or pentamethinecyanine
dyes I (Z1, 22 = atomic group forming benzene or naphthalene ring; Y1,

O, S, Se, NR; R = alkyl; R1, R2 = alkyl; R3-R5 = H, monovalent substituent; i = 0-2; R4 and R5 may form 5- or 6-member ring at i = 2; j

0, 1; the rings contain ≥1 thienyl or arylthio). The materials show low fog and high sensitivity. 208125-65-9
RL: MOA (Modifier or additive use); TEM (Technical or engineered material use); USES (Uses)
thermally developable silver halide photog. emulsion containing

sensitizer dye with low fog)
208125-65-9 CAPLUS
Benzothiazolium, 3-(5-carboxypentyl)-2-[5-[3-(5-carboxypentyl)-5-(2-thienyl)-2(3H)-benzothiazolylidene]-1,3-pentadienyl)-5-(2-thienyl)-,

inner salt (9CI) (CA INDEX NAME)

L7 ANSWER 31 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER:
1997:732508 CAPLUS
128:68551
Heat development photosensitive material useful in printing platemaking
INVENTOR(\$): Inagaki, Yoshior Oya, Toyohisa; Kobayashi, Katsu;
Tsuzuki, Hirohike
PATENT ASSIGNEE(\$): Fuji Photo Film Co., Ltd., Japan
SOURCE: Jpn. Kokai Tokkyo Koho, 22 pp.
CODEN: JKXXAF
DOCUMENT TYPE: Patent

DOCUMENT TYPE:

Patent Japanese

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 09292673	A	19971111	JP 1996-105788	19960425
JP 3606998	B2	20050105		
US 5948608	A	19990907	US 1997-840715	19970425
PRIORITY APPLN. INFO.:			JP 1996-105788 A	19960425

OTHER SOURCE(S): MARPAT 128:68551

The title material, possessing photosensitive Ag halide grains on ≥ 1 side of a support, contains a cyanine dye I $\{21,\ 22=$ atoms required to form benzene or naphthalene ring: Y1, Y2= O, S, Se, NR $\{R=alky1\}$: R1, R2= alky1: R3-9= H or substituent, R4 and R6, R5 and R7,

and R6 and R8 may link to form a 5 or 6-membered ring, R4, R6, and R8 may

link to form a ring in which 2 6-membered rings are condensed; X = counter ion;

i=0 or 1; R1-9, R, Z1 or Z2 has ≥ 1 thienyl or arylthic group as a substituent]. The material containing a binder, an organic Ag salt, a

reducing
agent for the salt, photosensitive Ag halide grains, may contain the dye
on >1 of a support. The material shows high sensitivity, low fog,
high contrast, and good storage stability before and after processing.

IT 200401-09-8
RL: DEV (Device component use); MOA (Modifier or additive use); USES
(Uses)
(heat)
(heat)
heptamethinecyanine
dye)
RN 200401-09-8 CAPLUS
CN Benzothiazolium, 3-(5-carboxypentyl)-2-[7-[3-(5-carboxypentyl)-5-[2-

L7 ANSWER 30 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

ANSWER 31 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued) thienyl)-2(3H)-benzothiazolylidene]-1,3,5-heptatrienyl]-5-(2-thienyl)-, inner salt (9CI) (CA INDEX NAME)

PAGE 1-B

127:169015
Photographic material containing emulsion sensitized with cyanine dye
Kagawa, Nobuski; Nakamura, Masaki; Ishii, Fumio
Konica Co., Japan
Jpn. Kokai Tokkyo Koho, 32 pp.
CODEN: JKXXAF

INVENTOR(S):
PATENT ASSIGNEE(S):
SOURCE:

DOCUMENT TYPE: Patent

LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 09179233	A	19970711	JP 1995-340159	19951227
JP 3333984 PRIORITY APPLN. INFO.:	B2	20021015	JP 1995-340159	19951227

GI

$$\begin{array}{c} v^{1} & v^{1} & v^{1} \\ v^{1} & (CL^{11})_{m11} & v^{12} \\ v^{2} & R^{11} & 0 \\ \end{array}$$

A photog, material has ≥ 1 emulsion layer sensitized with ≥ 1 cyanine dye of the structure I [21-2 = 5-6-membered N-heterocyclyl; Y1-2

NR, O, S, Se, Te; R = aliphatic group, aryl, heterocyclyl; R1-2 = C≤10 aliphatic group; R, R1 and/or R2 = group substituted with water-solubilizing group; W1 = H, F, lower aliphatic group, aryl; W2 = H, F; W1 and /or W2 = F;

L1-3 = (un) substituted methine; 11-2, m1-2 = 0, 1; m1 and /or m2 = 1; M1 number necessary for cancelling a total charge of the mol.; nl = number necessary for neutralizing a total charge of the mol.] or II [Y11-14 =

ANSWER 32 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued) NR10, O, S, Se, Te; R10 = aliph. group, aryl, heterocyclyl; R11-12 = CS10 aliph. group; R10, R11 and/or R12 = group substituted with water-solubilizing group; V1-4 = H, alkyl, alkoxy, aryl, V1 and V2, V3

V4 may be bound to each other to form a condensed ring; W11 = H. F. lower aliph, group, aryl; W12 = H. F.; W11 and/or W112 = F. L11-13 = (un) substituted methine; m11-12 = 0, 1; m11 and /or m12 = 1; M11 = no. necessary for cancelling a total charge of the mol.; n11 = no. necessary for meutralizing a total charge of the mol.]. The photog. material has high sensitivity and exhibits a small residual color after processing. R1: DEV (Device component unit)

193602-66-3
RI: DEV (Device component use); USES (Uses)
(photog. material containing emulsion sensitized with cyanine dye)
193602-66-3 CAPLUS
Thieno[3,2-d]thiazolium, 1-(carboxymethyl)-2-[{3-(carboxymethyl)-5-[2-[5-

(2-furanyl)-3-(3-sulfopropyl)-2(3H)-benzoxazolylidene]-1-methylethylidene]-4-oxo-2-thiazolidinylidene]fluoromethyl]-5,6-dihydro-, inner salt (9CI) (CA INDEX NAME)

ANSWER 33 OF 38 CAPLUS COPYRIGHT 2007 ACS ON STN SSION NUMBER: 1996:666522 CAPLUS MENT NUMBER: 125:288709

ACCESSION NUMBER: DOCUMENT NUMBER:

TITLE:

125:288709
Silver halide photographic material spectrally sensitized by trinuclear cyanine having improved red sensitivity and low dye stain Kagawa, Nobuaki; Kita, Noryasu Konishiroku Photo Ind., Japan Jpn. Kokai Tokkyo Koho, 30 pp. CODEN: XXXXAF Patent Japanese 1

INVENTOR(S):
PATENT ASSIGNEE(S):
SOURCE:

DOCUMENT TYPE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. DATE APPLICATION NO. KIND DATE JP 08201954 PRIORITY APPLN. INFO.: JP 1995-11332 JP 1995-11332 19950127 19950127 19960809

For diagram(s), see printed CA Issue. The claimed photog. material is characterized by (1) that ≥ 1 of the emulsion layer is spectrally sensitized by a cyanine dye Γ (21, 22=5-6).

6-membered heterocyclic ring; Z3 = NR, O, S, Se, Te; R, R2 = aliphatic,

ary1,
heterocyclic group; R1, R3 = C 1-10 aliphatic; at least one of R and
R1-3 has
a water-solubilizing group; L1 = substituted methine; L2, L3 = methyne;

and n = counter ion for stoichiometric balance; l, k, m = 0, l). A sensitizing dye II (Y11-13 = NR10, O, S, Se, Te; R10-13, L11-13 have the same meaning as R, R1-3, L1-3 in I; V1-4 = H, alkyl, aryl, alkoxy; ≥1 R10-13 has a water-solubilizing group; M1 and n = counter ion for stoichiometric balance; m = 0, l). The spectral sensitizer provides high sensitivity at red spectral region, and also provides the material with good shelf life and low residual dye stain at the processing. 182946-31-2 RL: DEV (Device component use); USES (Uses) (Ag halide photog, material spectrally sensitized by trinuclear ine

ine
having improved red sensitivity and low dye stain)
182946-31-2 CAPLUS
Thieno[3,2-d]thiazolium, 1-(carboxymethyl)-2-[[3-(carboxymethyl)-5-[2-[5-(2-furany)]-3-(3-sulfopropyl)-2(3H)-benzoxazolylidene]-2methoxyethylidene]-4-oxo-2-thiazolidinylidene]methyl]-, inner salt,

with N,N-diethylethanamine (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 182946-30-1 CMF C30 H25 N3 O11 S4

ANSWER 33 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

2

L7 ANSWER 34 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 1996:262402 CAPLUS
DOCUMENT NUMBER: 125:22191
Silver halide photographic material containing sensitizing methyne dye
Ooya, Toyohisa
PATENT ASSIGNEE(S): Fuji Photo Film Co Ltd, Japan
SOURCE: JPN Kokai Tokkyo Koho, 24 pp.
CODEN: JPCKAF
DOCUMENT TYPE: Patent
Japanese

Japanese

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE JP 1994-175430 JP 1994-175430 JP 08043984 А 19960216 19940727 PRIORITY APPLN. INFO.: 19940727

$$z^{1}$$
 z^{1}
 z^{1}
 z^{1}
 z^{1}
 z^{1}
 z^{1}
 z^{1}
 z^{1}
 z^{1}

The photog, material contains in ≥ 1 component layer a methyne compound I [XI = [substituted] group forming (condensed) 5- or 6-membered heterocycle: L1, L2 = [substituted] methyne; n = 0, 1; Q = (substituted) polymethyne or methyne; R1 = [substituted] aromatic or aliphatic; Z = ; I. =

connecting group; Ar = (substitute) aromatic; m = 0, 1; W1 = counter

The photog. material shows high sensitivity and low residual color stain. 177323-96-5
RL: DEV (Device component use); USES (Uses)
(Ag halide photog. material containing sensitizing methyne dye)
177323-96-5 CAPLUS
Benzolc acid, 2-{2-{5-(2-furanyl)-3-(4-sulfobutyl)-2(3H)-benzothiazolylidene]-1-[4-oxo-3-(phenylmethyl)-2-thioxo-5-thiazolidinylidene]-1, monopotassium salt (9CI) (CA INDEX NAME)

L7 ANSWER 35 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER:
1590:138971 CAPLUS
112:138971 CA

DOCUMENT TYPE: LANGUAGE: GI

3 New thiosemicarbazides I (R = Et, Bu Ph) were prepared from N-[4-phenyl-5-(2-thienyl)-1,2,4-triazol-3-yl]mercaptoacetohydrazide. Reaction of I with the appropriate phenacyl bromide afforded thiazolidederivs. II (R2 = H, Cl, Br, OMe) whereas their reaction with chloroacetic acid or maleic anhydride gave the corresponding thiazolidine derivs. Cyclodesulfurization of I with DCCD in PhMe yielded 5-substituted-amino-1,3,4-chiadiazoles. 6 Representative compds. were tested for their in-vitro antimicrobial activity against some pathogenic microorganisms, some of them were proved to be active. 125866-95-7P 125866-96-8P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of) 125866-35-7 CAPUUS 5-Thiazolidineacetic acid, 3-ethyl-4-oxo-2-[[[[4-phenyl-5-[2-thienyl]-4H-5-[3-thienyl]-4H-5-[3-thiazolidineacetic acid, 3-ethyl-4-oxo-2-[[[[4-phenyl-5-[2-thienyl]-4H-5-[3-thiazolidineacetic acid, 3-ethyl-4-oxo-2-[[[[4-phenyl-5-[2-thienyl]-4H-5-[3-thiazolidineacetic acid, 3-ethyl-4-oxo-2-[[[[4-phenyl-5-[2-thienyl]-4H-5-[3-thiazolidineacetic acid, 3-ethyl-4-oxo-2-[[[[4-phenyl-5-[2-thienyl]-4H-5-[3-thiazolidineacetic acid, 3-ethyl-4-oxo-2-[[[4-phenyl-5-[2-thienyl]-4H-5-[3-thiazolidineacetic acid, 3-ethyl-4-oxo-2-[[[4-phenyl-5-[2-thienyl]-4H-5-[3-thiazolidineacetic acid, 3-ethyl-4-oxo-2-[[[4-thienyl]-5-[3-thiazolidineacetic acid, 3-ethyl-4-oxo-2-[[[4-thienyl]-5-[3-thiazolidineacetic acid, 3-ethyl-4-oxo-2-[[[4-thienyl]-5-[3-thiazolidineacetic acid, 3-thyl-4-oxo-2-[[[4-thienyl]-5-[3-thiazolidineacetic acid, 3-thyl-4-oxo-2-[[[4-thienyl]-5-[3-thiazolidineacetic acid, 3-thyl-4-oxo-2-[[4-thienyl]-5-[3-thiazolidineacetic acid, 3-thyl-4-oxo-2-[[4-thienyl]-5-[4-thienyl]-4H-5-[4-

5-Thiazolidineacetic acid, 3-ethyl-4-oxo-2-[[[4-phenyl-5-(2-thienyl)-4H-1,2,4-triazol-3-yl]thio]acetyl]hydrazono]- (9CI) (CA INDEX NAME)

125866-96-8 CAPLUS

L7 ANSWER 34 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

ANSWER 35 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued 5-Thiazolidineacetic acid, 0-3-phenyl-2-{[[[4-phenyl-5-(2-thienyl)-4H-1,2,4-triazol-3-yl]thio]acetyl]hydrazono]- (9CI) (CA INDEX NAME)

L7 ANSWER 36 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1989:183059 CAPLUS DOCUMENT NUMBER: 110:183059

DOCUMENT NUMBER:

TITLE: Photosensitive composition for electrophotographic

heat-mode optical recording Kato, Eiichi; Ishii, Kazuo Fuji Photo Film Co., Ltd., Japan Jpn. Kokai Tokkyo Koho, 15 pp. CODEN: JKXXAF Patent: Japanese 1 INVENTOR(S): PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 63155146	A	19880628	JP 1986-301690	19861219
PRIORITY APPLN. INFO.:			JP 1986-301690	19861219

GI

$$Q = \begin{array}{c} 2 \\ \text{N} \\ \text{N} \\ \text{R} \end{array}$$

Electrophotog. photoreceptors and heat-mode optical recording medium use

atoms

to complete 5-membered heterocycle; R, R1 = aliphatic group; X, Y = 0, e:
A- = anion; r = 1, 2]. The sensitizer dye serves to improve the storage stability of electrophotog. photoreceptors and to improve the sensitivity of optical recording media at ≥750 nm.
120114-60-5
RE: USES (Uses)
(sensitizer dye, electrophotog. photoconductors and optical recording media from)

IT media from) 120114-60-5 CAPLUS

120114-60-5 CAPLUS

Benzothiazolium, 3-(3-carboxypropyl)-2-[5-[[5-[3-(3-carboxypropyl)-5-chloro-6-methoxy-2(3H)-benzothiazolylidene]-2(5H)-furanylidene]methyl]-2-furanyl]-5-chloro-6-methoxy-, tetrafluoroborate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 120114-59-2 CMF C33 H29 C12 N2 O8 S2

ANSWER 36 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)

CM 2

CRN 14874-70-5 CMF B F4 CCI CCS

L7 ANSWER 37 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 1962:58374 CAPLUS
DOCUMENT NUMBER: 56:58374
ORIGINAL REFERENCE NO.: 56:11113;,11114a-g
Sensitization of silver halide emulsions
INVENTOR(S): Bach, Guenther
DOCUMENT TYPE: Patent
LANGINGE: German INTERIOR (S):
INVENTOR(S):
DOCUMENT TYPE:
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DD 19683		19600818	DD	19570827
PRIORITY APPLN. INFO.:			DD	19570827

For diagram(s), see printed CA Issue. Cyanine and styryl dyes with 1 or more heterocyclic radicals of the general formula I, where R is a substituted heterocyclic radical, R' is

general formula I, where R is a substituted heterocyclic radical, R! is an alkyl, aralkyl, or aryl radical, X is O, S, Se, NR", CH:CR", or C(Y)2 radical (Y and Z are awl, alkyl or cycloalkyl groups), R" is H, alkyl, cycloalkyl, or aryl radicals, are prepared AgCl emulsion (1 kg.) containing 40

mg. N,N'-diethylquinopseudocyanine was sensitized with 2 mg.
5-(2-benzoxazolyl)-2(4-dimethylaminostyryl)benzothiazole (II), sensitization range 500-590 mm, maximum 580 mm. II. m. 246-8' (C6H6) is obtained by melting p-Me2NC6H4CHO with 2-methyl-5- (2benzoxazolyl)benzothiazole (III) and anhydrous zncl2 for 3 hrs.
2-(4-Chloro-3-nitrophenyl)benzoxazole (IV), m. 178-80' (BuoH), was treated with Na252 and then with a mixture of AcOH-Ac20-Zn in dioxane at 120-40' to give III, m. 158-80' (MeoH), Melting 1 g.
2-mechyl-3-(2-furyl)benzoxazole (V) with 0 tg. ethylene sulfate at 110', condensation with 8 ml. EtC(Obt.) (V) in 5 ml. pyridine and 3 ml. AcOH for 2 hrs. at 150', and precipitation with Et20 gave 9-ethyl5,5'-di-2-furyl-3,3'-bis(2-sulfatoethyl)oxacarbocyanine inner salt,

m. 226° (decomposition) (MeOH), sensitization range 500-580 m μ , maximum 560-65 m μ . V, m. 60-2° (petr. ether), b5-6 154-5°, was obtained by boiling 2-methyl-5-nitroso-acetamidobenzoxazole (VII) with furan for 80 hrs. and vacuum distillation of the residue after removal

furan for 80 hrs. and vacuum distillation of the residue alone.

of the
furan excess. 2-Methyl-5-(2-thienyl)benzoxazole, m. 68-70° (EtOH),
which is obtained from VII and thiophene (VIII), was melted with 2.6 g.
Et 2,4-dinitrobenzenesulfonate; 2.5 g. of the quaternary salt was heated
with 1.2 g. S-ethylisothiopropionanilide for 30 min. at 180°. Of
this product, 1.5 g. was condensed with 0.95 g. 2-methyl-3-ethyl-5methoxybenzothiazolium p-toluenesulfonate in 3 ml. pyridineAcOH (5:1)
mixture (IX) for 1 hr. at 125°. The dye is precipitated with 2 ml. 20%
acueous

aqueous

KI and extracted with H2O to give

3-ethyl-2-[2-ethyl-3-(3-ethyl-5-methoxy-2-benzothiazolinylidene)propenyl]-5-(2-thienyl)benzoxazolium iodide, m.p.
203-5*, sensitization range 500-620 mµ, maximum 600-5. Heating

1.15 g. 2-methyl-6-(2-thienyl)benzothiazole (X) with 0.76 g. Br(CH2)2CO2H
for 1 hr. and boiling with 2 ml. VI in 10 ml. IX gave

3,3'-bis(2-carboxyethyl)-5-ethyl-6,6'-di-2-thienylthiacarbocyanine inner
salt, m. 204-6' (EtOH), sensitization range 500-650 mµ. X, m.
83-5' (EtOH), was obtained by diazotizing 2-methyl-6aminobenzothiazole, stirring the diazonium salt solution with VIII and 5N

ANSWER 37 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued) NaOH for 15 hrs. Heating 1.6 g. 1-phenyl-2-methyl-5-(2-benzoxazolyl)benzimidazole (XI) with 5 ml. MeI in a sealed tube for 3

benzowazolyl)benzimidazole (XI) with 5 ml. Mei in a sealed tube for 3 at 100°, boiling the product with 1 ml. HC(OEt)3 in 10 ml. PhNO2, and pptn. with Et20 gave 5;5'-di-2benzowazolyl-3,3'-dimethyl-1,1'-diphenylazacyanine iodide m. 270° (Me2CO), sensitization max. 560-5 mμ. XI, m. 127-8' (petr. ether), is prepd. by boiling IV with PNNH2, redn. of 2-(4-anilino-3-nitrophenyl)benzoxazole, m. 161-3' (Me2CO) with H in MeOH and Raney Ni as catalyst, and boiling the diamine with AcOH-Ac2O.

105730-01-6P, 3-(2-carboxyethyl)-2-[2-[[3-(2-carboxyethyl)-6-(2-thienyl)-2-benzothiazolinylidene]methyl]-1-butenyl]-6-(2-thienyl)-2-(2-flamignum manufacture of, and use as sensitizer in photography)

105730-01-6 CAPLUS

3-(2-Carboxyethyl)-2-[2-[[3-(2-carboxyethyl)-6-(2-thienyl)-2-benzothiazolinylidene]methyl]-1-butenyl]-6-(2-thienyl)benzothiazolium hydroxide, inner salt (6CI, 7CI) (CA INDEX NAME)

L7 ANSWER 38 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 1961:92058 CAPLUS
OCIUMENT NUMBER: 55:92058
ORIGINAL REFERENCE NO: 55:17321f-1,17322a-b
SITILE: Sensitization of silver halide emulsions
INVENTOR(s): Bach, Gunther
VEB Filmfabrik Agfa Wolfen
DOCUMENT TYPE: Pater
LANGUAGE: Pater
Unavailable
FMHILY ACCL NUM. COUNT: 1

TITLE:
INVENTOR(S):
PATENT ASSIGNEE(S):
DOCUMENT TYPE:
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:

PATENT NO. APPLICATION NO. KIND DATE DE 1063028 1957-v13077 19570910
Analogs of 5,5'-diphenyl-3,3',9-triethylbenzoxacarbocyanine carrying a bulky heterocyclic substituent provided increased sensitivity in the 500-90 mµ range without increased sensitivity to red, while maintaining spectral selectivity and absence of undesirable diffusion. Thus, Purple Coupler z 169 (Cornwell-Clyne, Colour Cinematography, 1951, p. 384 (CA

47,
3735f)) and 3.0 mg. of each sensitizer were mixed with 100 cc. AgBr-AgI emulsion and cast on a film base. Test strips were illuminated by a spectrograph to equal exposures and developed black-and-white in a solution containing hydroquinone and p-methylaminophenol to determine the range of sensitization. The dyes were prepared by heating a substituted 2-methyl N-heterocyclic compound, optionally quaternized, with an ortho ester or derivative, suitably in a mixture of AcOH and pyridine at 125-40°. For example, 2-(4-chloro-3-nitrophenyl]benzoxazole (I), m. 178-80°, and NaZSZ gave the disulfide; reductive cyclization with Zn, AcOH, and AcOO in

dioxane at 120-40° gave 2-methyl-5-(2-benzoxazolyl)benzothiazole, m. 158-60°, which on fusion with p-Me2NC6H6CHO and ZnC12 for 3 hrs. gave a dye, m. 246-8°; sensitivity maximum 580 mµ, range 500-90 mµ. 2-Methyl-5-(2-furyl)benzoxazole, b. 154-5°, m. 60-2°, prepared from furan and 2-methyl-5-(N-mitrosoacetamido)benzoxazole, was quaternized with ethylene sulfate and condensed with EtC(OEt)3 (II) to give a dye, m. 226°, with sensitivity maximum 560-5 mµ, range 500-80 mµ. 2-Methyl-5-(2-thienyl)benzoxazole, m. 68-70°, was quaternized with 2,4-(NO2)2C6H4SO3Et, heated for 30 min. at 180° with EtC(SEt):NHPh, and the resulting intermediate was condensed with 2-methyl-3-ethyl-5-methoxybenzothiazolium salt; the dye was precipitated with KI and tallized from

methoxybenzothiazolium salt; the dye was precipitated which is also crystallized from alc., m. 203-5°, sensitivity maximum 600-5 mµ, range 500-620 mµ. Diazotized 2-methyl-6-aminobenzothiazole, thiophene, and 5N NaOH gave the 6-(2-thienyl)lderivative m. 83-5°; this was quaternized with BrCHZCHZCO2H and condensed with II to give a dye, m. 204-6°, sensitivity range 500-650 mµ. I and boiling PhNHZ gave the anilino derivative, m. 161-3°; this was reduced with H and Raney Ni in MeOH and the diamine boiled with AcOH-Ac2O to give 1-phenyl-2-methyl-5-(2-benzoxazolyl)benzimidazole, m. 127-8°, which was quaternized with MeI, condensed with HC(OEt)3 in PhNO2, and precipitated with ether to give a dye,

a dye, m. 270°, sensitivity maximum 560-5 m μ . Cf. CA 53, 4985c.

ANSWER 38 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN (Continued) 105730-01-6P, 3-{2-Carboxyethyl}-2-[2-{[3-{2-carboxyethyl}-6-{2-thienyl}-benzothiazolinylidene|methyl]-1-butenyl]-6-{2-thienyl}-benzothiazolium hydroxide, inner salt RL: PREP (Preparation of) (preparation of) 105730-01-6 CAPLUS 3-{2-Carboxyethyl}-2-[2-{[3-{2-carboxyethyl}-6-{2-thienyl}-2-benzothiazolinylidene|methyl}-1-butenyl]-6-{2-thienyl}-benzothiazolium hydroxide, inner salt (6CI, 7CI) (CA INDEX NAME)